Abstract

The Dakota toolkit provides a flexible and extensible interface between simulation codes and iterative analysis methods. Dakota contains algorithms for optimization with gradient and nongradient-based methods; uncertainty quantification with sampling, reliability, and stochastic expansion methods; parameter estimation with nonlinear least squares methods; and sensitivity/variance analysis with design of experiments and parameter study methods. These capabilities may be used on their own or as components within advanced strategies such as surrogate-based optimization, mixed integer nonlinear programming, or optimization under uncertainty. By employing object-oriented design to implement abstractions of the key components required for iterative systems analyses, the Dakota toolkit provides a flexible and extensible problem-solving environment for design and performance analysis of computational models on high performance computers.

This report serves as a reference manual for the commands specification for the Dakota software, providing input overviews, option descriptions, and example specifications.
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The Dakota software (http://dakota.sandia.gov/) delivers advanced parametric analysis techniques enabling quantification of margins and uncertainty, risk analysis, model calibration, and design exploration with computational models. Its methods include optimization, uncertainty quantification, parameter estimation, and sensitivity analysis, which may be used individually or as components within surrogate-based and other advanced strategies.

Author


The Reference Manual documents all the input keywords that can appear in a Dakota input file to configure a Dakota study. Its organization closely mirrors the structure of dakota.input.summary. For more information see Dakota Input Specification. For information on software structure, refer to the Developers Manual [3], and for a tour of Dakota features and capabilities, including a tutorial, refer to the User's Manual[5].

Dakota 6.14 and newer include a refreshed Examples Library, ranging from examples of Dakota input files and basic studies through complete case studies. These can be found in binary distributions in dakota/share/dakota/examples/official (dakota/dakota-examples in source distributions). Legacy examples, including many from the Dakota software manuals appear in other directories in dakota/share/dakota/examples (dakota/examples for source).

1.1 How to Use this Manual

- To learn how to run Dakota from the command line, see Running Dakota
- To learn how to restart Dakota studies, see Restarting Dakota Studies
- To learn about the Dakota restart utility, see The Dakota Restart Utility

To find more information about a specific keyword

1. Use the search box at the top right (currently only finds keyword names)
2. Browse the Keywords tree on the left navigation pane
3. Look at the Dakota Input Specification
4. Navigate through the keyword pages, starting from the Keywords Area

To find more information about a Dakota related topic
1. Browse the Topics Area on the left navigation pane
2. Navigate through the topics pages, starting from the Topics Area

A small number of examples are included (see Sample Input Files) along with a description of the test problems (see Test Problems).

A bibliography for the Reference Manual is provided in Bibliographic References
Chapter 2

Running Dakota

The Dakota executable file is named dakota (dakota.exe on Windows) and is most commonly run from a terminal or command prompt.

2.1 Usage

If the dakota command is entered at the command prompt without any arguments, a usage message similar to the following appears:

usage: dakota [options and <args>]
      -help (Print this summary)
      -version (Print DAKOTA version number)
      -input <$val> (REQUIRED DAKOTA input file $val)
      -preproc [$val] (Pre-process input file with pyprepro or tool $val)
      -output <$val> (Redirect DAKOTA standard output to file $val)
      -error <$val> (Redirect DAKOTA standard error to file $val)
      -parser <$val> (Parsing technology: nidr[strict][:dumpfile])
      -no_input_echo (Do not echo DAKOTA input file)
      -check (Perform input checks)
      -pre_run [$val] (Perform pre-run (variables generation) phase)
      -run [$val] (Perform run (model evaluation) phase)
      -post_run [$val] (Perform post-run (final results) phase)
      -read_restart [$val] (Read an existing DAKOTA restart file $val)
      -stop_restart <$val> (Stop restart file processing at evaluation $val)
      -write_restart [$val] (Write a new DAKOTA restart file $val)

Of these command line options, only input is required, and the -input switch can be omitted if the input file name is the final item appearing on the command line (see Examples); all other command-line inputs are optional.

- help prints the usage message above.
- version prints version information for the executable.
- check invokes a dry-run mode in which the input file is processed and checked for errors, but the study is not performed.
- input provides the name of the Dakota input file, which can optionally be pre-processed as a template using the preproc option.
- output and error options provide file names for redirection of the Dakota standard output (stdout) and standard error (stderr), respectively.
- The parser option is for debugging and will not be further described here.
• By default, Dakota will echo the input file to the output stream, but `no_input_echo` can override this behavior.

• `read_restart` and `write_restart` commands provide the names of restart databases to read from and write to, respectively.

• `stop_restart` command limits the number of function evaluations read from the restart database (the default is all the evaluations) for those cases in which some evaluations were erroneous or corrupted. Restart management is an important technique for retaining data from expensive engineering applications.

• `-pre_run`, `-run`, and `-post_run` instruct Dakota to run one or more execution phases, excluding others. The commands must be followed by filenames as described in Execution Phases.

Command line switches can be abbreviated so long as the abbreviation is unique, so the following are valid, unambiguous specifications: `-h`, `-v`, `-c`, `-i`, `-o`, `-e`, `-s`, `-w`, `-re`, `-ru`, and `-po` and can be used in place of the longer forms of the command line options.

For information on restarting Dakota, see Restarting Dakota Studies and The Dakota Restart Utility.

2.2 Examples

To run Dakota with a particular input file, the following syntax can be used:

    dakota -i dakota.in

or more simply

    dakota dakota.in

This will echo the standard output (stdout) and standard error (stderr) messages to the terminal. To redirect stdout and stderr to separate files, the `-o` and `-e` command line options may be used:

    dakota -i dakota.in -o dakota.out -e dakota.err

or

    dakota -o dakota.out -e dakota.err dakota.in

Alternatively, any of a variety of Unix redirection variants can be used. Refer to[6] for more information on Unix redirection. The simplest of these redirects stdout to another file:

    dakota dakota.in > dakota.out

The specified Dakota input file may instead be a dprepro/aprepro-style template file to be pre-processed prior to running Dakota. For example it might contain template expressions in curly braces:

```
# {MyLB = 2.0} {MyUB = 8.6}
variables
    uniform_uncertain 3
    upper_bounds {MyUB} {MyUB} {MyUB}
    lower_bounds {MyLB} {MyLB} {MyLB}
```

(See the Interfaces chapter in the Dakota User's Manual[?] for more information and use cases.) To pre-process the input file, specify the `preproc` flag which generates an intermediate temporary input file for use in Dakota. If Dakota's `pyprepro.py` utility is not available on the execution PATH and/or additional pre-processing options are needed, the tool location and syntax can be specified, for example:

---

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2.3 Execution Phases

Dakota has three execution phases: pre-run, run, and post-run.

• **pre-run** can be used to generate variable sets

• **run** (core run) invokes the simulation to evaluate variables, producing responses

• **post-run** accepts variable/response sets and analyzes the results (for example, calculate correlations from a set of samples). Currently only two modes are supported and only for sampling, parameter study, and DACE methods:

1. **pre-run** only with optional tabular output of variables:
   
   ```
   dakota -i dakota.in -pre_run [::myvariables.dat]
   ```

2. **post-run** only with required tabular input of variables/responses:
   
   ```
   dakota -i dakota.in -post_run myvarsresponses.dat::
   ```

2.4 Restarting Dakota Studies

Dakota is often used to solve problems that require repeatedly running computationally expensive simulation codes. In some cases you may want to repeat an optimization study, but with a tighter final convergence tolerance. This would be costly if the entire optimization analysis had to be repeated. Interruptions imposed by computer usage policies, power outages, and system failures could also result in costly delays. However, Dakota automatically records the variable and response data from all function evaluations so that new executions of Dakota can pick up where previous executions left off. The Dakota restart file (``dakota.rst``` by default) archives the tabulated interface evaluations in a binary format. The primary restart commands at the command line are **-read_restart**, **-write_restart**, and **-stop_restart**.

2.4.1 Writing Restart Files

To write a restart file using a particular name, the **-write_restart** command line input (may be abbreviated as **-w**) is used:

```
    dakota -i dakota.in -write_restart my_restart_file
```

If no **-write_restart** specification is used, then Dakota will still write a restart file, but using the default name `dakota.rst` instead of a user-specified name.

To turn restart recording off, the user may use the **restart_file** keyword, in the interface block. This can increase execution speed and reduce disk storage requirements, but at the expense of a loss in the ability to recover and continue a run that terminates prematurely. This option is not recommended when function evaluations are costly or prone to failure. Please note that using the **deactivate restart_file** specification will result in a zero length restart file with the default name `dakota.rst`, which can overwrite an exiting file.
2.4.2 Using Restart Files

To restart Dakota from a restart file, the `-read_restart` command line input (may be abbreviated as `-r`) is used:

```
dakota -i dakota.in -read_restart my_restart_file
```

If no `-read_restart` specification is used, then Dakota will not read restart information from any file (i.e., the default is no restart processing).

To read in only a portion of a restart file, the `-stop_restart` control (may be abbreviated as `-s`) is used to specify the number of entries to be read from the database. Note that this integer value corresponds to the restart record processing counter (as can be seen when using the `print` utility (see The Dakota Restart Utility) which may differ from the evaluation numbers used in the previous run if, for example, any duplicates were detected (since these duplicates are not recorded in the restart file). In the case of a `-stop_restart` specification, it is usually desirable to specify a new restart file using `-write_restart` so as to remove the records of erroneous or corrupted function evaluations. For example, to read in the first 50 evaluations from `dakota.rst`:

```
dakota -i dakota.in -r dakota.rst -s 50 -w dakota_new.rst
```

The `dakota_new.rst` file will contain the 50 processed evaluations from `dakota.rst` as well as any new evaluations. All evaluations following the 50th in `dakota.rst` have been removed from the latest restart record.

2.4.3 Appending to a Restart File

If the `-write_restart` and `-read_restart` specifications identify the same file (including the case where `-write_restart` is not specified and `-read_restart` identifies `dakota.rst`), then new evaluations will be appended to the existing restart file.

2.4.4 Working with multiple Restart Files

If the `-write_restart` and `-read_restart` specifications identify different files, then the evaluations read from the file identified by `-read_restart` are first written to the `-write_restart` file. Any new evaluations are then appended to the `-write_restart` file. In this way, restart operations can be chained together indefinitely with the assurance that all of the relevant evaluations are present in the latest restart file.

2.4.5 How it Works

Dakota’s restart algorithm relies on its duplicate detection capabilities. Processing a restart file populates the list of function evaluations that have been performed. Then, when the study is restarted, it is started from the beginning (not a warm start) and many of the function evaluations requested by the iterator are intercepted by the duplicate detection code. This approach has the primary advantage of restoring the complete state of the iteration (including the ability to correctly detect subsequent duplicates) for all methods/iterators without the need for iterator-specific restart code. However, the possibility exists for numerical round-off error to cause a divergence between the evaluations performed in the previous and restarted studies. This has been rare in practice.

2.5 The Dakota Restart Utility

The Dakota restart utility program provides a variety of facilities for managing restart files from Dakota executions. The executable program name is `dakota_restart_util` and it has the following options, as shown by the usage message returned when executing the utility without any options:

```
Usage:
  dakota_restart_util command <arg1> [<arg2> <arg3> ...] --options
dakota_restart_util print <restart_file>
dakota_restart_util to_neutral <restart_file> <neutral_file>
dakota_restart_util from_neutral <neutral_file> <restart_file>
dakota_restart_util to_tabular <restart_file> <text_file>
```
2.5 The Dakota Restart Utility

Several of these functions involve format conversions. In particular, the binary format used for restart files can be converted to ASCII text and printed to the screen, converted to and from a neutral file format, or converted to a tabular format for importing into third-party plotting programs. In addition, a restart file with corrupted data can be repaired by value or id, and multiple restart files can be combined to create a master database.

2.5.1 Print Command

The `print` option is useful to show contents of a restart file, since the binary format is not convenient for direct inspection. The restart data is printed in full precision, so that exact matching of points is possible for restarted runs or corrupted data removals. For example, the following command

dakota_restart_util print
dakota.rst

results in output similar to the following:

```
-------------------------------
Restart record 1 (evaluation id 1):
-------------------------------
Parameters:
  1.8000000000000000e+00 intake_dia
  1.0000000000000000e+00 flatness

Active response data:
Active set vector = [ 3 3 3 ]
-2.4355973813420619e+00 obj_fn
-4.7428486677140930e-01 nln_ineq_con_1
-4.5000000000000000e-01 nln_ineq_con_2
  1.3971143170299741e-01 nln_ineq_con_3
[
  -4.364429863447897e-01 1.4999999999999999e-01 ] obj_fn gradient
  [ 2.9814239699997572e+01 0.0000000000000000e+00 ] nln_ineq_con_1 gradient
  [ 0.0000000000000000e+00 1.4999999999999999e-01 ] nln_ineq_con_2 gradient
  [ 0.0000000000000000e+00 -1.9485571585149869e-01 ] nln_ineq_con_3 gradient
-------------------------------
Restart record 2 (evaluation id 2):
-------------------------------
Parameters:
  2.1640000000000000e+00 intake_dia
  1.7169994018008317e+00 flatness

Active response data:
Active set vector = [ 3 3 3 ]
-2.4869127192988878e+00 obj_fn
  6.9256987999999843e-01 nln_ineq_con_1
  3.4245008972987528e-01 nln_ineq_con_2
  8.7142207937157910e-03 nln_ineq_con_3
[
  -4.364429863447897e-01 1.4999999999999999e-01 ] obj_fn gradient
  [ 2.9814239699997572e+01 0.0000000000000000e+00 ] nln_ineq_con_1 gradient
  [ 0.0000000000000000e+00 1.4999999999999999e-01 ] nln_ineq_con_2 gradient
  [ 0.0000000000000000e+00 -1.6998301774282701e-01 ] nln_ineq_con_3 gradient
...
```

Restart file processing completed: 11 evaluations retrieved.
2.5.2 Neutral File Format

A Dakota restart file can be converted to a neutral file format using a command like the following:

```
dakota_restart_util to_neutral dakota.rst dakota.neu
```

which results in a report similar to the following:

```
Writing neutral file dakota.neu
Restart file processing completed: 11 evaluations retrieved.
```

Similarly, a neutral file can be returned to binary format using a command like the following:

```
dakota_restart_util from_neutral dakota.neu dakota.rst
```

which results in a report similar to the following:

```
Reading neutral file dakota.neu
Writing new restart file dakota.rst
Neutral file processing completed: 11 evaluations retrieved.
```

The contents of the generated neutral file are similar to the following (from the first two records for the Cylinder example in[5]).

```
6 7 2 1.8000000000000000e+00 intake_dia 1.0000000000000000e+00 flatness 0 0 0 0 0
NULL 4 2 1 0 3 3 3 1 2 obj_fn nln_ineg_con_1 nln_ineg_con_2 nln_ineg_con_3
-2.4355973813420619e+00 -4.742848677140930e-01 -4.5000000000000001e-01
1.3971143170299741e-01 -4.364298963447897e-01 1.4999999999999999e-01
1.3855136437818300e-01 0.0000000000000000e+00 0.0000000000000000e+00
1.4999999999999999e-01 0.0000000000000000e+00 -1.9485571585149869e-01
6 7 2 2.1640000000000001e+00 intake_dia 1.7169994018008317e+00 flatness 0 0 0 0 0
NULL 4 2 1 0 3 3 3 1 2 obj_fn nln_ineg_con_1 nln_ineg_con_2 nln_ineg_con_3
-2.4869127192988878e+00 6.9256958799989843e-01 -3.4245008972987528e-01
8.7142207937157910e-03 -4.364298963447897e-01 1.4999999999999999e-01
2.981423969997572e+01 0.0000000000000000e+00 0.0000000000000000e+00
1.4999999999999999e-01 0.0000000000000000e+00 -1.6998301774287201e-01
```

This format is not intended for direct viewing (print should be used for this purpose). Rather, the neutral file capability has been used in the past for managing portability of restart data across platforms (recent use of more portable binary formats has largely eliminated this need) or for advanced repair of restart records (in cases where the remove command was insufficient).

2.5.3 Tabular Format

Conversion of a binary restart file to a tabular format enables convenient import of this data into 3rd-party post-processing tools such as Matlab, TECplot, Excel, etc. This facility is nearly identical to the output activated by the tabular_data keyword in the Dakota input file specification, but with two important differences:

1. No function evaluations are suppressed as they are with tabular_data (i.e., any internal finite difference evaluations are included).
2. The conversion can be performed later, i.e., for Dakota runs executed previously.

An example command for converting a restart file to tabular format is:

```
dakota_restart_util to_tabular dakota.rst dakota.m
```

which results in a report similar to the following:

```
Writing tabular text file dakota.m
Restart file processing completed: 10 evaluations tabulated.
```
2.5 The Dakota Restart Utility

The contents of the generated tabular file are similar to the following (from the example in the Restart section of [5]). Note that while evaluations resulting from numerical derivative offsets would be reported (as described above), derivatives returned as part of the evaluations are not reported (since they do not readily fit within a compact tabular format):

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
4 NO_ID 0.9 1.10011 0.0002004407265 0.259945 0.7602420121
5 NO_ID 0.9 1.09989 0.0001995607255 0.260055 0.7597580121
6 NO_ID 0.58256179 0.4772244441 0.1050555937 0.1007670171 -0.06353963386
7 NO_ID 0.5826200462 0.4772244441 0.1050386469 0.1008348962 -0.06356876195
8 NO_ID 0.5825035339 0.4772244441 0.1050725476 0.1006991449 -0.06351050577
9 NO_ID 0.58256179 0.4772701663 0.1050283245 0.100743156 -0.06349408333
10 NO_ID 0.58256179 0.4771747219 0.1050828704 0.1007908783 -0.06358517983
...```

Controlling tabular format: The command-line options --freeform and --custom_annotated give control of headers in the resulting tabular file. Freeform will generate a tabular file with no leading row nor columns (variable and response values only). Custom annotated format accepts any or all of the options:

- **header:** include %-commented header row with labels
- **eval_id:** include leading column with evaluation ID
- **interface_id:** include leading column with interface ID

For example, to recover Dakota 6.0 tabular format, which contained a header row, leading column with evaluation ID, but no interface ID:

```
dakota_restart_util to_tabular dakota.rst dakota.m --custom_annotated header eval_id
```

Resulting in

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Finally, **--output_precision integer** will generate tabular output with the specified integer digits of precision.

2.5.4 Concatenation of Multiple Restart Files

In some instances, it is useful to combine restart files into a single master function evaluation database. For example, when constructing a data fit surrogate model, data from previous studies can be pulled in and reused to create a combined data set for the surrogate fit. An example command for concatenating multiple restart files is:

```
dakota_restart_util cat dakota.rst.1 dakota.rst.2 dakota.rst.3 dakota.rst.all
```

which results in a report similar to the following:

```
Writing new restart file dakota.rst.all
dakota.rst.1 processing completed: 10 evaluations retrieved.
dakota.rst.2 processing completed: 110 evaluations retrieved.
dakota.rst.3 processing completed: 65 evaluations retrieved.
```

The dakota.rst.all database now contains 185 evaluations and can be read in for use in a subsequent Dakota study using the **-read_restart** option to the dakota executable.
2.5.5 Removal of Corrupted Data

On occasion, a simulation or computer system failure may cause a corruption of the Dakota restart file. For example, a simulation crash may result in failure of a post-processor to retrieve meaningful data. If 0's (or other erroneous data) are returned from the user’s analysis driver, then this bad data will get recorded in the restart file. If there is a clear demarcation of where corruption initiated (typical in a process with feedback, such as gradient-based optimization), then use of the -stop_restart option for the dakota executable can be effective in continuing the study from the point immediately prior to the introduction of bad data. If, however, there are interspersed corruptions throughout the restart database (typical in a process without feedback, such as sampling), then the remove and remove_ids options of dakota_restart_util can be useful.

An example of the command syntax for the remove option is:

```bash
dakota_restart_util remove 2.e-04 dakota.rst dakota.rst.repaired
```

which results in a report similar to the following:

```bash
Writing new restart file dakota.rst.repaired
Restart repair completed: 65 evaluations retrieved, 2 removed, 63 saved.
```

where any evaluations in dakota.rst having an active response function value that matches 2.e-04 within machine precision are discarded when creating dakota.rst.repaired.

An example of the command syntax for the remove_ids option is:

```bash
dakota_restart_util remove_ids 12 15 23 44 57 dakota.rst dakota.rst.repaired
```

which results in a report similar to the following:

```bash
Writing new restart file dakota.rst.repaired
Restart repair completed: 65 evaluations retrieved, 5 removed, 60 saved.
```

where evaluation ids 12, 15, 23, 44, and 57 have been discarded when creating dakota.rst.repaired.

An important detail is that, unlike the -stop_restart option which operates on restart record numbers, the remove_ids option operates on evaluation ids. Thus, removal is not necessarily based on the order of appearance in the restart file. This distinction is important when removing restart records for a run that contained either asynchronous or duplicate evaluations, since the restart insertion order and evaluation ids may not correspond in these cases (asynchronous evaluations have ids assigned in the order of job creation but are inserted in the restart file in the order of job completion, and duplicate evaluations are not recorded which introduces offsets between evaluation id and record number). This can also be important if removing records from a concatenated restart file, since the same evaluation id could appear more than once. In this case, all evaluation records with ids matching the remove_ids list will be removed.

If neither of these removal options is sufficient to handle a particular restart repair need, then the fallback position is to resort to direct editing of a neutral file to perform the necessary modifications. <!-----------------------------

------>
Chapter 3

Dakota HDF5 Output

Beginning with release 6.9, Dakota gained the ability to write many method results such as the correlation matrices computed by sampling studies and the best parameters discovered by optimization methods to disk in HDF5. In Dakota 6.10 and above, evaluation data (variables and responses for each model or interface evaluation) may also be written. Many users may find this newly supported format more convenient than scraping or copying and pasting from Dakota’s console output.

To enable HDF5 output, the results_output keyword with the hdf5 option must be added to the Dakota input file. In addition, Dakota must have been built with HDF5 support. Beginning with Dakota 6.10, HDF5 is enabled in our publicly available downloads. HDF5 support is considered a somewhat experimental feature. The results of some Dakota methods are not yet written to HDF5, and in a few, limited situations, enabling HDF5 will cause Dakota to crash.

3.1 HDF5 Concepts

HDF5 is a format that is widely used in scientific software for efficiently storing and organizing data. The HDF5 standard and libraries are maintained by the HDF Group.

In HDF5, data are stored in multidimensional arrays called datasets. Datasets are organized hierarchically in groups, which also can contain other groups. Datasets and groups are conceptually similar to files and directories in a filesystem. In fact, every HDF5 file contains at least one group, the root group, denoted “/”, and groups and datasets are referred to using slash-delimited absolute or relative paths, which are more accurately called link names.

HDF5 has as one goal that data be “self-documenting” through the use of metadata. Dakota output files include two kinds of metadata.

- **Dimension Scales.** Each dimension of a dataset may have zero or more scales, which are themselves datasets. Scales are often used to provide, for example, labels analogous to column headings in a table (see the dimension scales that Dakota applies to moments) or numerical values of an independent variable (user-specified probability levels in level mappings).

- **Attributes.** key:value pairs that annotate a group or dataset. A key is always a character string, such as dakota_version, and (in Dakota output) the value can be a string-, integer-, or real-valued scalar. Dakota stores the number of samples that were requested in a sampling study in the attribute ‘samples’.

3.2 Accessing Results

Many popular programming languages have support, either natively or from a third-party library, for reading and writing HDF5 files. The HDF Group itself supports C/C++ and Java libraries. The Dakota Project suggests the h5py module for Python. Examples that demonstrate using h5py to access and use Dakota HDF5 output may be found in the Dakota installation at dakota/share/dakota/examples/official/hdf5.
3.3 Organization of Results

Currently, complete or nearly complete coverage of results from sampling, optimization and calibration methods, parameter studies, and stochastic expansions exists. Coverage will continue to expand in future releases to include not only the results of all methods, but other potentially useful information such as interface evaluations and model transformations.

Methods in Dakota have a character string Id and are executed by Dakota one or more times. (Methods are executed more than once in studies that include a nested model, for example.) The Id may be provided by the user in the input file using the id_method keyword, or it may be automatically generated by Dakota. Dakota uses the label NO_METHOD_ID for methods that are specified in the input file without an id_method, and NOSPEC_METHOD_ID_<N> for methods that it generates for its own internal use. The <N> in the latter case is an incrementing integer that begins at 1.

The results for the <N>th execution of a method that has the label <method Id> are stored in the group

```
/methods/<method Id>/results/execution:<N>/
```

The /methods group is always present in Dakota HDF5 files, provided at least one method added results to the output. (In a future Dakota release, the top level groups /interfaces and /models will be added.) The group execution:1 is also always present, even if there is only a single execution.

The groups and datasets for each type of result that Dakota is currently capable of storing are described in the following sections. Every dataset is documented in its own table. These tables include:

- A brief description of the dataset.
- The location of the dataset relative to /methods/<method Id>/execution:<N>. This path may include both literal text that is always present and replacement text. Replacement text is <enclosed in angle brackets and italicized>. Two examples of replacement text are <response descriptor> and <variable descriptor>, which indicate that the name of a Dakota response or variable makes up a portion of the path.
- Clarifying notes, where appropriate.
- The type (String, Integer, or Real) of the information in the dataset.
- The shape of the dataset; that is, the number of dimensions and the size of each dimension.
- A description of the dataset’s scales, which includes
  - The dimension of the dataset that the scale belongs to.
  - The type (String, Integer, or Real) of the information in the scale.
  - The label or name of the scale.
  - The contents of the scale. Contents that appear in plaintext are literal and will always be present in a scale. Italicized text describes content that varies.
  - Notes that provide further clarification about the scale.
- A description of the dataset’s attributes, which are key:value pairs that provide helpful context for the dataset.

The Expected Output section of each method’s keyword documentation indicates the kinds of output, if any, that method currently can write to HDF5. These are typically in the form of bulleted lists with clarifying notes that refer back to the sections that follow.

3.3.1 Study Metadata

Several pieces of information about the Dakota study are stored as attributes of the top-level HDF5 root group ("/"). These include:
3.3 Organization of Results

3.3.2 A Note about Variables Storage

Variables in most Dakota output (e.g. tabular data files) and input (e.g. imported data to construct surrogates) are listed in “input spec” order. (The variables keyword section is arranged by input spec order.) In this ordering, they are sorted first by function:

1. Design
2. Aleatory
3. Epistemic
4. State

And within each of these categories, they are sorted by domain:

1. Continuous
2. Discrete integer (sets and ranges)
3. Discrete string
4. Discrete real

A shortcoming of HDF5 is that datasets are homogeneous; for example, string- and real-valued data cannot readily be stored in the same dataset. As a result, Dakota has chosen to flip “input spec” order for HDF5 and sort first by domain, then by function when storing variable information. When applicable, there may be as many as four datasets to store variable information: one to store continuous variables, another to store discrete integer variables, and so on. Within each of these, variables will be ordered by function.

3.3.3 Sampling Moments

Sampling produces moments (e.g. mean, standard deviation or variance) of all responses, as well as 95% lower and upper confidence intervals for the 1st and 2nd moments. These are stored as described below. When sampling is used in incremental mode by specifying refinement_samples, all results, including the moments group, are placed within groups named increment:<N>, where <N> indicates the increment number beginning with 1.

3.3.4 Correlations

A few different methods produce information about the correlations between pairs of variables and responses (collectively: factors). The four tables in this section describe how correlation information is stored. One important note is that HDF5 has no special, native type for symmetric matrices, and so the simple correlations and simple rank correlations are stored in dense 2D datasets.

3.3.5 Probability Density

Some aleatory UQ methods estimate the probability density of responses.

3.3.6 Level Mappings

Aleatory UQ methods can calculate level mappings (from user-specified probability, reliability, or generalized reliability to response, or vice versa).
3.3.7 Variance-Based Decomposition (Sobol' Indices)

Dakota’s sampling method can produce main and total effects; stochastic expansions (polynomial_choas, stoch_-collocation) additionally can produce interaction effects.

Each order (pair, 3-way, 4-way, etc) of interaction is stored in a separate dataset. The scales are unusual in that they are two-dimensional to contain the labels of the variables that participate in each interaction.

3.3.8 Integration and Expansion Moments

Stochastic expansion methods can obtain moments two ways.

3.3.9 Extreme Responses

Sampling with epistemic variables produces extreme values (minimum and maximum) for each response.

3.3.10 Parameter Sets

All parameter studies (vector_parameter_study, list_parameter_study, multidim_parameter_study, centered_-parameter_study) record tables of evaluations (parameter-response pairs), similar to Dakota’s tabular output file. Centered parameter studies additionally store evaluations in an order that is more natural to interpret, which is described below.

In the tabular-like listing, variables are stored according to the scheme described in a previous section.

3.3.11 Variable Slices

Centered parameter studies store “slices” of the tabular data that make evaluating the effects of each variable on each response more convenient. The steps for each individual variable, including the initial or center point, and corresponding responses are stored in separate groups.

3.3.12 Best Parameters

Dakota’s optimization and calibration methods report the parameters at the best point (or points, for multiple final solutions) discovered. These are stored using the scheme described in the variables section. When more than one solution is reported, the best parameters are nested in groups named set:<N>, where <N> is a integer numbering the set and beginning with 1.

State (and other inactive variables) are reported when using objective functions and for some calibration studies. However, when using configuration variables in a calibration, state variables are suppressed.

3.3.13 Best Objective Functions

Dakota’s optimization methods report the objective functions at the best point (or points, for multiple final solutions) discovered. When more than one solution is reported, the best objective functions are nested in groups named set:<N>, where <N> is a integer numbering the set and beginning with 1.

3.3.14 Best Nonlinear Constraints

Dakota’s optimization and calibration methods report the nonlinear constraints at the best point (or points, for multiple final solutions) discovered. When more than one solution is reported, the best constraints are nested in groups named set:<N>, where N is a integer numbering the set and beginning with 1.
3.3.15 Calibration

When using calibration terms with an optimization method, or when using a nonlinear least squares method such as nl2sol, Dakota reports residuals and residual norms for the best point (or points, for multiple final solutions) discovered.

3.3.16 Parameter Confidence Intervals

Least squares methods (nl2sol, nlssol_sqp, optpp_g_newton) compute confidence intervals on the calibration parameters.

3.3.17 Best Model Responses (without configuration variables)

When performing calibration with experimental data (but no configuration variables), Dakota records, in addition to the best residuals, the best original model responses.

3.3.18 Best Model Responses (with configuration variables)

When performing calibration with experimental data that includes configuration variables, Dakota reports the best model responses for each experiment. These results include the configuration variables, stored in the scheme described in the variables section, and the model responses.

3.3.19 Multistart and Pareto Set

The multi_start and pareto_set methods are meta-iterators that control multiple optimization sub-iterators. For both methods, Dakota stores the results of the sub-iterators (best parameters and best results). For multi_start, Dakota additionally stores the initial points, and for pareto_set, it stores the objective function weights.

3.4 Organization of Evaluations

An evaluation is a mapping from variables to responses performed by a Dakota model or interface. Beginning with release 6.10, Dakota has the ability to report evaluation history in HDF5 format. The HDF5 format offers many advantages over existing console output and tabular output. Requiring no “scraping”, it is more convenient for most users than the former, and being unrestricted to a two-dimensional, tabular arrangement of information, it is far richer than the latter.

This section begins by describing the Dakota components that can generate evaluation data. It then documents the high-level organization of the data from those components. Detailed documentation of the individual datasets (the “low-level” organization) where data are stored follows. Finally, information is provided concerning input keywords that control which components report evaluations.

3.4.1 Sources of Evaluation Data

Evaluation data are produced by only two kinds of components in Dakota: models and interfaces. The purpose of this subsection is to provide a basic description of models and interfaces for the purpose of equipping users to manage and understand HDF5-format evaluation data.

Because interfaces and models must be specified in even simple Dakota studies, most novice users of Dakota will have some familiarity with these concepts. However, the exact nature of the relationship between methods, models, and interfaces may be unclear. Moreover, the models and interfaces present in a Dakota study are not always limited to those specified by the user. Some input keywords or combinations of components cause Dakota to create new models or interfaces “behind the scenes” and without the user’s direct knowledge. Not only can user-specified models and interfaces write evaluation data to HDF5, but also these auto-generated components. Accordingly, it
may be helpful for consumers of Dakota’s evaluation data to have a basic understanding of how Dakota creates and employs models and interfaces.

Consider first the input file shown here.

```plaintext
environment
tabular_data
results_output
   hdf5

method
   id_method 'sampling'
sampling
   samples 20
   model_pointer 'sim'

model
   id_model 'sim'
   single
   interface_pointer 'tb'

variables
   uniform_uncertain 2
   descriptors 'x1' 'x2'
   lower_bounds 0.0 0.0
   upper_bounds 1.0 1.0

responses
   response_functions 1
   descriptors 'f'
   no_gradients
   no_hessians

interface
   id_interface 'tb'
   fork
   analysis_drivers 'text_book'
```

This simple input file specifies a single method of type sampling, which also has the Id ‘sampling’. The ‘sampling’ method possesses a model of type single (alias simulation) named ‘sim’, which it uses to perform evaluations. (Dakota would have automatically generated a single model had one not been specified.) That is to say, for each variables-to-response mapping required by the method, it provides variables to the model and receives back responses from it.

Single/simulation models like ‘sim’ perform evaluations by means of an interface, typically an interface to an external simulation. In this case, the interface is ‘tb’. The model passes the variables to ‘tb’, which executes the text_book driver, and receives back responses.

It is clear that two components produce evaluation data in this study. The first is the single model ‘sim’, which receives and fulfills evaluation requests from the method ‘sampling’, and the second is the interface ‘tb’, which similarly receives requests from ‘sim’ and fulfills them by running the text_book driver.

Because tabular data was requested in the environment block, a record of the model’s evaluations will be reported to a tabular file. The interface’s evaluations could be dumped from the restart file using dakota_restart_util.

If we compared these evaluation histories from ‘sim’ and ‘tb’, we would see that they are identical to one another. The model ‘sim’ is a mere “middle man” whose only responsibility is passing variables from the method down to the interface, executing the interface, and passing responses back up to the method. However, this is not always the case.

For example, if this study were converted to a gradient-based optimization using optpp_q_newton, and the user specified numerical_gradients:

```plaintext
# model and interface same as above. Replace the method, variables, and responses with:

method
   id_method 'opt'
optpp_q_newton

variables
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Then the model would have the responsibility of performing finite differencing to estimate gradients of the response \( f \) requested by the method. Multiple function evaluations of ‘tb’ would map to a single gradient evaluation at the model level, and the evaluation histories of ‘sim’ and ‘tb’ would contain different information.

Note that because it is unwieldy to report gradients (or Hessians) in a tabular format, they are not written to the tabular file, and historically were available only in the console output. The HDF5 format provides convenient access to both the “raw” evaluations performed by the interface and higher level model evaluations that include estimated gradients.

This pair of examples hopefully provides a basic understanding of the flow of evaluation data between a method, model, and interface, and explains why models and interfaces are producers of evaluation data.

Next consider a somewhat more complex study that includes a Dakota model of type surrogate. A surrogate model performs evaluations requested by a method by executing a special kind of interface called an approximation interface, which Dakota implicitly creates without the direct knowledge of the user. Approximation interfaces are a generic container for the various kinds of surrogates Dakota can use, such as gaussian processes.

A Dakota model of type global surrogate may use a user-specified dace method to construct the actual underlying model(s) that it evaluates via its approximation interface. The dace method will have its own model (typically of type single/simulation), which will have a user-specified interface.

In this more complicated case there are at least four components that produce evaluation data: (1) the surrogate model and (2) its approximation interface, and (3) the dace method’s model and (4) its interface. Although only components (1), (3), and (4) are user-specified, evaluation data produced by (2) may be written to HDF5, as well. (As explained below, only evaluations performed by the surrogate model and the dace interface will be recorded by default. This can be overridden using hdf5 sub-keywords.) This is an example where “extra” and potentially confusing data appears in Dakota’s output due to an auto-generated component.

An important family of implicitly-created models is the recast models, which have the responsibility of transforming variables and responses. One type of recast called a data transform model is responsible for computing residuals when a user provides experimental data in a calibration study. Scaling recast models are employed when scaling is requested by the user for variables and/or responses.

Recast models work on the principle of function composition, and “wrap” a submodel, which may itself also be a recast model. The innermost model in the recursion often will be the simulation or surrogate model specified by the user in the input file. Dakota is capable of recording evaluation data at each level of recast.

### 3.4.2 High-level Organization of Evaluation Data

This subsection describes how evaluation data produced by models and interfaces are organized at high level. A detailed description of the datasets and subgroups that contain evaluation data for a specific model or interface is given in the next subsection.

Two top level groups contain evaluation data, /interfaces and /models.

#### 3.4.2.1 Interfaces

Because interfaces can be executed by more than one model, interface evaluations are more precisely thought of as evaluations of an interface/model combination. Consequently, interface evaluations are grouped not only by interface Id (‘tb’ in the example above), but also the Id of the model that requested them (‘sim’).
If the user does not provide an Id for an interface that he specifies, Dakota assigns it the Id NO_ID. Approximation interfaces receive the Id APPROX_INTERFACE_<N>, where N is an incrementing integer beginning at 1. Other kinds of automatically generated interfaces are named NOSPEC_INTERFACE_ID_<N>.

3.4.2.2 Models

The top-level group for model evaluations is /models. Within this group, model evaluations are grouped by type: simulation, surrogate, nested, or recast, and then by model Id. That is:

/mdels/<type>/<model Id>/

Similar to interfaces, user-specified models that lack an Id are given one by Dakota. A single model is named NO_MODEL_ID. Some automatically generated models receive the name NOSPEC_MODEL_ID.

Recast models are a special case and receive the name RECAST_<WRAPPED-MODEL>_TYPE_<N>. In this string:

- **WRAPPED-MODEL** is the Id of the innermost wrapped model, typically a user-specified model
- **TYPE** is the specific kind of recast. The three most common recasts are:
  - RECAST: several generic responsibilities, including summing objective functions to present to a single-objective optimizer
  - DATA_TRANSFORM: Compute residuals in a calibration
  - SCALING: scale variables and responses
- N is an incrementing integer that begins with 1. It is employed to distinguish recasts of the same type that wrap the same underlying model.

The model’s evaluations may be the result of combining information from multiple sources. A simulation/single model will receive all the information it requires from its interface, but more complicated model types may use information not only from interfaces, but also other models and the results of method executions. Nested models, for instance, receive information from a submethod (the mean of a response from a sampling study, for instance) and potentially also an optional interface.

The sources of a model’s evaluations may be roughly identified by examining the contents of that models’ sources group. The sources group contains softlinks (note: softlinks are an HDF5 feature analogous to soft or symbolic links on many file systems) to groups for the interfaces, models, or methods that the model used to produce its evaluation data. (At this time, Dakota does not report the specific interface or model evaluations or method executions that were used to produce a specific model evaluation, but this is a planned feature.)

Method results likewise have a sources group that identifies the models or methods employed by that method. By following the softlinks contained in a method’s or model’s sources group, it is possible to “drill down” from a method to its ultimate sources of information. In the sampling example above, interface evaluations performed via the ‘sim’ model at the request of the ‘sampling’ method could be obtained at the HDF5 path: /methods/sampling/sources/sim/sources/tb/

3.4.3 Low-Level Organization of Evaluation Data

The evaluation data for each interface and model are stored using the same schema in a collection of groups and datasets that reside within that interface or model’s high-level location in the HDF5 file. This section describes that “low-level” schema.

Data are divided first of all into variables, responses, and metadata groups.

3.4.3.1 Variables

The variables group contains datasets that store the variables information for each evaluation. Four datasets may be present, one for each “domain”: continuous, discrete_integer, discrete_string, and
3.4 Organization of Evaluations

Discrete data. These datasets are two-dimensional, with a row (0th dimension) for each evaluation and a column (1st dimension) for each variable. The 0th dimension has one dimension scale for the integer-valued evaluation Id. The 1st dimension has two scales. The 0th scale contains descriptors of the variables, and the 1st contains their variable Ids. In this context, the Ids are a 1-to-N ranking of the variables in Dakota “input spec” order.

3.4.3.2 Responses

The responses group contains datasets for functions and, when available, gradients and Hessians.

**Functions:** The functions dataset is two-dimensional and contains function values for all responses. Like the variables datasets, evaluations are stored along the 0th dimension, and responses are stored along the 1st. The evaluation Ids and response descriptors are attached as scales to these axes, respectively.

**Gradients:** The gradients dataset is three-dimensional. It has the shape evaluations \( \times \) responses \( \times \) variables. Dakota supports a specification of mixed_gradients, and the gradients dataset is sized and organized such that only those responses for which gradients are available are stored. When mixed_gradients are employed, a response will not necessarily have the same index in the functions and gradients datasets.

Because it is possible that the gradient could be computed with respect to any of the continuous variables, active or inactive, that belong to the associated model, the gradients dataset is sized to accommodate gradients taken with respect to all continuous variables. Components that were not included in a particular evaluation will be set to NaN (not a number), and the derivative_variables_vector (in the metadata group) for that evaluation can be examined as well.

**Hessians:** Hessians are stored in a four-dimensional dataset, evaluations \( \times \) responses \( \times \) variables \( \times \) variables. The hessians dataset shares many of the characteristics with the gradients: in the mixed_hessians case, it will be smaller in the response dimension than the functions dataset, and unrequested components are set to NaN.

3.4.3.3 Metadata

The metadata group contains up to three datasets.

**Active Set Vector:** The first is the active_set_vector. It is two dimensional, with rows corresponding to evaluations and columns corresponding to responses. Each element contains an integer in the range 0-7, which indicates the request (function, gradient, Hessian) for the corresponding response for that evaluation. The 0th dimension has the evaluations Ids scale, and the 1st dimension has two scales: the response descriptors and the "default" or "maximal" ASV, an integer 0-7 for each response that indicates the information (function, gradient, Hessian) that possibly could have been requested during the study.

**Derivative Variables Vector:** The second dataset in the metadata group is the derivative_variables_vector dataset. It is included only when gradients or Hessians are available. Like the ASV, it is two-dimensional. Each column of the DVV dataset corresponds to a continuous variable and contains a 0 or 1, indicating whether gradients and Hessians were computed with respect to that variable for the evaluation. The 0th dimension has the evaluation Ids as a scale, and the 1st dimension has two scales. The 0th is the descriptors of the continuous variables. The 1st contains the variable Ids of the continuous variables.

**Analysis Components:** The final dataset in the metadata group is the analysis_components dataset. It is a 1D dataset that is present only when the user specified analysis components, and it contains those components as strings.

3.4.4 Selecting Models and Interfaces to Store

When HDF5 output is enabled (by including the hdf5 keyword), then by default evaluation data for the following components will be stored:

- The model that belongs to the top-level method. (Currently, if the top-level method is a metaiterator such as method-hybrid, no model evaluation data will be stored.)
- All simulation interfaces. (interfaces of type fork, system, direct, etc.)
The user can override these defaults using the keywords `model_selection` and `interface_selection.

The choices for `model_selection` are:

- **top_method**: (default) Store evaluation data for the top method's model only.
- **all_methods**: Store evaluation data for all models that belong directly to a method. Note that these models may be recasts of user-specified models, not the user-specified models themselves.
- **all**: Store evaluation data for all models.
- **none**: Store evaluation data for no models.

The choices for `interface_selection` are:

- **simulation**: (default) Store evaluation data for simulation interfaces.
- **all**: Store evaluation data for all interfaces.
- **none**: Store evaluation data for no interfaces.

If a model or interface is excluded from storage by these selections, then they cannot appear in the `sources` group for methods or models.

### 3.5 Distribution Parameters

Variables are characterized by parameters such as the mean and standard deviation or lower and upper bounds. Typically, users provide these parameters as part of their input to Dakota, but Dakota itself may also compute them as it scales and transforms variables, normalizes empirical distributions (e.g. for `histogram_bin_uncertain` variables), or calculates alternative parameterizations (lambda and zeta vs mean and standard deviation for a `lognormal_uncertain`).

Beginning with release 6.11, models write their variable's parameters to HDF5. The information is located in each model's `metadata/variable_parameters` subgroup. Within this group, parameters are stored by Dakota variable type (e.g. `normal_uncertain`), with one 1D dataset per type. The datasets have the same names as their variable types and have one element per variable. Parameters are stored by name.

Consider the following variable specification, which includes two normal and two uniform variables:

```
variables
  normal_uncertain 2
  descriptors 'nuv_1' 'nuv_2'
  means 0.0 1.0
  std deviations 1.0 0.5
  uniform_uncertain 2
  descriptors 'uuv_1' 'uuv_2'
  lower bounds -1.0 0.0
  upper bounds 1.0 1.0
```

Given this specification, and assuming a model ID of "tb_model", Dakota will write two 1D datasets, both of length 2, to the group `/models/simulation/tb_model/metadata/variable_parameters`, the first named `normal_uncertain`, and the second named `uniform_uncertain`. Using a JSON-like representation for illustration, the `normal_uncertain` dataset will appear as:

```
[
  {
    "mean": 0.0,
    "std_deviation": 1.0,
    "lower_bound": -inf,
    "upper_bound": inf
  },
  {
    "mean": 1.0,
```
3.5 Distribution Parameters

The uniform_uncertain dataset will contain:

```python
[{
    "lower_bound": -1.0,
    "upper_bound": 1.0
},
{
    "lower_bound": 0.0,
    "upper_bound": 1.0
}]
```

In these representations of the normal_uncertain and uniform_uncertain datasets, the outer square brackets ([]) enclose the dataset, and each element within the datasets are enclosed in curly braces ({}). The curly braces are meant to indicate that the elements are dictionary-like objects that support access by string field name.

A bit more concretely, the following code snippet demonstrates reading the mean of the second normal variable, nuv_2.

```python
import h5py
with h5py.File("dakota_results.h5") as h:
    model = h["/models/simulation/tb_model/"]
    # nu_vars is the dataset that contains distribution parameters for
    # normal_uncertain variables
    nu_vars = model["variable_parameters/normal_uncertain"]
    nuv_2_mu = nu_vars[1]["mean"] # 1 is the 0-based index of nuv_2, and
    # "mean" is the name of the field where
    # the mean is stored; nuv_2_mu now contains
    # 1.0.
```

The feature in HDF5 that underlies this name-based storage of fields is compound datatypes, which are similar to C/C++ structs or Python dictionaries. Further information about how to work with compound datatypes is available in the h5py documentation.

3.5.1 Naming Conventions and Layout

In most cases, datasets for storing parameters have names that match their variable types. The normal_uncertain and uniform_uncertain datasets illustrated above are examples. Exceptions include types such as discrete_design_set, which has string, integer, and real subtypes. For these, the dataset name is the top-level type with _string, _int, or _real appended: discrete_design_set_string, discrete_design_set_int, and discrete_design_set_real.

Most Dakota variable types have scalar parameters. For these, the names of the parameters are generally the singular form of the associated Dakota keyword. For example, triangular_uncertain variables are characterized in Dakota input using the plural keywords modes, lower_bounds, and upper_bounds. The singular field names are, respectively, "mode", "lower_bound", and "upper_bound". In this case, all three parameters are real-valued and stored as floating point numbers, but variable types/fields can also be integer-valued (e.g. binomial_uncertain/num_trials) or string-valued.

Some variable/parameter fields contain 1D arrays or vectors of information. Consider histogram_bin_uncertain variables, for which the user specifies not just one value, but an ordered collection of abscissas and corresponding ordinates or counts. Dakota stores the abscissas in the "abscissas" field, which is a 1D dataset of floating-point numbers. It similarly stores the counts in the "counts" field. (In this case, only the normalized counts are stored, regardless of whether the user provided counts or ordinates.)

When the user specifies more than one histogram_bin_uncertain variable, it often is also necessary to include the pairs_per_variable keyword to divide the abscissa/count pairs among the variables. This raises the question of how lists of parameters that vary in length across the variables ought to be stored.
Although HDF5 supports variable-length datasets, for simplicity (and due to limitations in h5py at the time of the 6.11 release), Dakota stores vector parameter fields in conventional fixed-length datasets. The lengths of these datasets are determined at runtime in the following way: For a particular variable type and field, the field for all variables is sized to be large enough to accommodate the variable with the longest list of parameters. Any unused space for a particular variable is filled with NaN (if the parameter is real-valued), INTMAX (integer-valued), or an empty string (string-valued). In addition, each variable has an additional field, “num_elements”, that reports the number of elements in the fields that contain actual data and not fill values.

Consider this example, in which the user has specified a pair of histogram_bin_uncertain variables. The first has 3 pairs, and the second has 4.

variables
  histogram_bin_uncertain 2
  pairs_per_variable 2 3
  abscissas 0.0 0.5 1.0
  -1.0 -0.5 0.5 1.0
  counts 0.25 0.75 0.0
  0.2 0.4 0.2 0.0

For this specification, Dakota will write a dataset named histogram_bin_uncertain to the metadata/variable_parameters/ subgroup for the model. It will be of length 2, one element for each variable, and contain the following:

```
[
  {
    "num_elements": 3,
    "abscissas": [0.0, 0.5, 1.0, NaN],
    "counts": [0.25, 0.75, 0.0, NaN]
  },
  {
    "num_elements": 4,
    "abscissas": [-1.0, -0.5, 0.5, 1.0],
    "counts": [0.2, 0.4, 0.2, 0.0]
  }
]
```

3.5.2 h5py Examples

The fields available for a variable parameters dataset can be determined in h5py by examining the datatype of the dataset.

```
1 import h5py
2 with h5py.File("dakota_results.h5") as h:
3   model = h['/models/simulation/NO_MODEL_ID/']
4   md = model['metadata/variable_parameters']
5   nu = md['normal_uncertain']
6   nu_param_names = nu.dtype.names
7   # nu_param_names is a tuple of strings: ('mean', 'std_deviation',
8     # 'lower_bound', 'upper_bound')

3.5.3 Known Limitations

h5py has a known bug that prevents parameters for some types of variables from being accessed (the Python interpreter crashes with a segfault). These include:

- histogram_point_uncertain string
- discrete_uncertain_set string

3.5.4 Metadata

The variable parameter datasets have two dimension scales. The first (index 0) contains the variable descriptors, and the second (index 1) contains variable Ids. Available Parameters
3.5 Distribution Parameters

3.5.5 Parameter Listing for All Types

The table below lists all Dakota variables and parameters that can be stored.
Chapter 4

Test Problems

This page contains additional information about two test problems that are used in Dakota examples throughout the Dakota manuals *Textbook* and *Rosenbrock*.

Many of these examples are also used as code verification tests. The examples are run periodically and the results are checked against known solutions. This ensures that the algorithms are correctly implemented.

Additional test problems are described in the User’s Manual.

4.1 Textbook

The two-variable version of the “textbook” test problem provides a nonlinearly constrained optimization test case. It is formulated as:

\[
\begin{align*}
\text{minimize } f &= (x_1 - 1)^4 + (x_2 - 1)^4 \\
\text{subject to } g_1 &= x_1^2 - \frac{x_2}{2} \leq 0 \\
& \quad g_2 = x_2^2 - \frac{x_1}{2} \leq 0 \\
& \quad 0.5 \leq x_1 \leq 5.8 \\
& \quad -2.9 \leq x_2 \leq 2.9
\end{align*}
\]

Contours of this test problem are illustrated in the next two figures.

![Figure 4.1: Contours of the textbook problem on the [-3,4] x [-3,4] domain. The feasible region lies at the intersection of the two constraints g_1 (solid) and g_2 (dashed).](image-url)
For the textbook test problem, the unconstrained minimum occurs at \((x_1, x_2) = (1, 1)\). However, the inclusion of the constraints moves the minimum to \((x_1, x_2) = (0.5, 0.5)\). Equation \text{textbookform} presents the 2-dimensional form of the textbook problem. An extended formulation is stated as

\[
\begin{align*}
\text{minimize } f &= \sum_{i=1}^{n} (x_i - 1)^4 \\
\text{subject to } g_1 &= x_1^2 - \frac{x_2}{2} \leq 0 \\
g_2 &= x_2^2 - \frac{x_1}{2} \leq 0 \\
0.5 &\leq x_1 \leq 5.8 \\
-2.9 &\leq x_2 \leq 2.9
\end{align*}
\]

where \(n\) is the number of design variables. The objective function is designed to accommodate an arbitrary number of design variables in order to allow flexible testing of a variety of data sets. Contour plots for the \(n = 2\) case have been shown previously.

For the optimization problem given in Equation \text{tbe}, the unconstrained solution (\text{num\_nonlinear\_inequality\_constraints set to zero}) for two design variables is:

\[
\begin{align*}
x_1 &= 1.0 \\
x_2 &= 1.0
\end{align*}
\]

with

\[
f^* = 0.0
\]

The solution for the optimization problem constrained by \(g_1\) (\text{num\_nonlinear\_inequality\_constraints set to one}) is:

\[
\begin{align*}
x_1 &= 0.763 \\
x_2 &= 1.16
\end{align*}
\]

with

\[
\begin{align*}
f^* &= 0.00388 \\
g_1^* &= 0.0 \text{ (active)}
\end{align*}
\]
The solution for the optimization problem constrained by \( g_1 \) and \( g_2 \) (num_nonlinear_inequality_constraints set to two) is:

\[
\begin{align*}
    x_1 &= 0.500 \\
    x_2 &= 0.500
\end{align*}
\]

with

\[
\begin{align*}
    f^* &= 0.125 \\
    g_1^* &= 0.0 \text{ (active)} \\
    g_2^* &= 0.0 \text{ (active)}
\end{align*}
\]

Note that as constraints are added, the design freedom is restricted (the additional constraints are active at the solution) and an increase in the optimal objective function is observed.

### 4.2 Rosenbrock

The Rosenbrock function[34] is a well-known test problem for optimization algorithms. The standard formulation includes two design variables, and computes a single objective function. This problem can also be posed as a least-squares optimization problem with two residuals to be minimized because the objective function is the sum of squared terms.

#### Standard Formulation

The standard two-dimensional formulation can be stated as

\[
\text{minimize } f = 100(x_2-x_1^2)^2 + (1-x_1)^2 \tag{rosenstd}
\]

Surface and contour plots for this function are shown in the Dakota User’s Manual.

The optimal solution is:

\[
\begin{align*}
    x_1 &= 1.0 \\
    x_2 &= 1.0
\end{align*}
\]

with

\[
    f^* = 0.0
\]

#### A Least-Squares Optimization Formulation

This test problem may also be used to exercise least-squares solution methods by recasting the standard problem formulation into:

\[
\text{minimize } f = (f_1)^2 + (f_2)^2 \tag{rosenls}
\]

where

\[ f_1 = 10(x_2-x_1^2) \tag{rosenr1} \]

and

\[ f_2 = 1-x_1 \tag{rosenr2} \]

are residual terms.

The included analysis driver can handle both formulations. In the dakota/share/dakota/test directory, the rosenbrock executable (compiled from Dakota_Source/test/rosenbrock.cpp) checks the number of response functions passed in the parameters file and returns either an objective function (as computed from Equation rosenstd) for use with optimization methods or two least squares terms (as computed from Equations rosenr1 -rosenr2 ) for use with least squares methods. Both cases support analytic gradients of the function set with respect to the design variables. See the User’s Manual for examples of both cases (search for Rosenbrock).
Chapter 5

Dakota Input Specification

Dakota input is specified in a text file, e.g., `dakota_uq.in` containing blocks of keywords that control program behavior. This section describes the format and admissible elements of an input file.

5.1 Dakota Keywords

Valid Dakota input keywords are dictated by `dakota.xml`, included in source and binary distributions of Dakota. This specification file is used with the NIDR[30] parser to validate user input and is therefore the definitive source for input syntax, capability options, and optional and required capability sub-parameters for any given Dakota version. A more readable variant of the specification `dakota.input.summary` is also distributed.

While complete, users may find `dakota.input.summary` overwhelming or confusing and will likely derive more benefit from adapting example input files to a particular problem. Some examples can be found here: Sample Input Files. Advanced users can master the many input specification possibilities by understanding the structure of the input specification file.

5.2 Input Spec Overview

Refer to the `dakota.input.summary` file, in Input Spec Summary, for all current valid input keywords.

- The summary describes every keyword including:
  - Whether it is required or optional
  - Whether it takes ARGUMENTS (always required) Additional notes about ARGUMENTS can be found here: Specifying Arguments.
  - Whether it has an ALIAS, or synonym
  - Which additional keywords can be specified to change its behavior

- Additional details and descriptions are described in Keywords Area

- For additional details on NIDR specification logic and rules, refer to[30] (Gay, 2008).

5.2.1 Common Specification Mistakes

Spelling mistakes and omission of required parameters are the most common errors. Some causes of errors are more obscure:

- Documentation of new capability sometimes lags its availability in source and executables, especially stable releases. When parsing errors occur that the documentation cannot explain, reference to the particular input
specification `dakota.input.summary` used in building the executable, which is installed alongside the executable, will often resolve the errors.

- If you want to compare results with those obtained using an earlier version of Dakota (prior to 4.1), your input file for the earlier version must use backslashes to indicate continuation lines for Dakota keywords. For example, rather than

```plaintext
# Comment about the following "responses" keyword...
responses,
  objective_functions = 1
# Comment within keyword "responses"
analytic_gradients
# Another comment within keyword "responses"
no_hessians
```

you would need to write

```plaintext
# Comment about the following "responses" keyword...
responses, \
  objective_functions = 1 \ 
# Comment within keyword "responses" \ 
analytic_gradients \ 
# Another comment within keyword "responses" \ 
no_hessians
```

with no white space (blanks or tabs) after the `\` character.

In most cases, the Dakota parser provides error messages that help the user isolate errors in input files. Running `dakota -input dakota_study.in -check` will validate the input file without running the study.

### 5.2.2 Specifying Arguments

Some keywords, such as those providing bounds on variables, have an associated list of values or strings, referred to as arguments.

When the same value should be repeated several times in a row, you can use the notation `N*value` instead of repeating the value `N` times.

For example

```plaintext
lower_bounds -2.0 -2.0 -2.0
upper_bounds 2.0 2.0 2.0
```

could also be written

```plaintext
lower_bounds 3*-2.0
upper_bounds 3* 2.0
```

(with optional spaces around the `*`).

Another possible abbreviation is for sequences: `L:S:U` (with optional spaces around the `:`) is expanded to `L L+S L+2*S ... U`, and `L:U` (with no second colon) is treated as `L:1:U`.

For example, in one of the test examples distributed with Dakota (test case 2 of `test/dakota_uq_textbook_sop_lhs.in`),

```plaintext
histogram_point = 2
abscissas = 50. 60. 70. 80. 90.
            30. 40. 50. 60. 70.
counts = 10 20 30 20 10
         10 20 30 20 10
```

could also be written

```plaintext
histogram_point = 2
abscissas = 50 : 10 : 90
            30 : 10 : 70
counts = 10:10:30 20 10
        10:10:30 20 10
```
Count and sequence abbreviations can be used together. For example

```
response_levels =
    0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0
    0.0 0.1 0.2 0.3 0.4 0.5 0.6 0.7 0.8 0.9 1.0
```
can be abbreviated

```
response_levels =
    2*0.0:0.1:1.0
```

### 5.3 Sample Input Files

A Dakota input file is a collection of fields from the dakota.input.summary file that describe the problem to be solved by Dakota. Several brief examples follow below.

Dakota 6.14 and newer include a refreshed Examples Library, ranging from examples of Dakota input files and basic studies through complete case studies. These can be found in binary distributions in `dakota/share/dakota/examples/official` (dakota/dakota-examples in source distributions). Legacy examples, including many from the Dakota software manuals appear in other directories in `dakota/share/dakota/examples` (dakota/examples for source).

#### Sample 1: Optimization

The following sample input file shows single-method optimization of the Textbook Example (see Textbook) using DOT’s modified method of feasible directions. A similar file is available as `dakota/share/dakota/examples/users/textbook_opt_conmin.in`.

```
# Dakota Input File: textbook_opt_conmin.in
environment
    tabular_data
        tabular_data_file = 'textbook_opt_conmin.dat'
method
    # dot_mmfd #DOT performs better but may not be available
    conmin_mfd
        max_iterations = 50
        convergence_tolerance = 1e-4
variables
    continuous_design = 2
    initial_point 0.9 1.1
    upper_bounds 5.8 2.9
    lower_bounds 0.5 -2.9
    descriptors 'x1' 'x2'
interface
    direct
        analysis_driver = 'text_book'
responses
    objective_functions = 1
    nonlinear_inequality_constraints = 2
    numerical_gradients
        method_source dakota
        interval_type central
        fd_gradient_step_size = 1.e-4
    no_hessians
```

#### Sample 2: Least Squares (Calibration)

The following sample input file shows a nonlinear least squares (calibration) solution of the Rosenbrock Example (see Rosenbrock) using the NL2SOL method. A similar file is available as `dakota/share/dakota/examples/users/rosen_opt_nls.in`.

```
# Dakota Input File: rosen_opt_nls.in
```
environment
tabular_data
tabular_data_file = 'rosen_opt_nls.dat'

method
max_iterations = 100
convergence_tolerance = 1e-4
nl2sol

model
single

variables
continuous_design = 2
initial_point = -1.2 1.0
lower_bounds = -2.0 -2.0
upper_bounds = 2.0 2.0
descriptors = 'x1' 'x2'

interface
analysis_driver = 'rosenbrock'
direct

responses
calibration_terms = 2
analytic_gradients
no_hessians

Sample 3: Nondeterministic Analysis

The following sample input file shows Latin Hypercube Monte Carlo sampling using the Textbook Example (see Textbook). A similar file is available as dakota/share/dakota/test/dakota_uq_textbook_lhs.in.

method,
sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
sample_type lhs

variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'

histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 10 1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'

histogram_point_uncertain = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'

interface,
fork asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses,
response_functions = 3
Sample 4: Parameter Study

The following sample input file shows a 1-D vector parameter study using the Textbook Example (see Textbook). It makes use of the default environment and model specifications, so they can be omitted. A similar file is available in the test directory as dakota/share/dakota/examples/users/rosen_ps_vector.in.

```dakota
# Dakota Input File: rosen_ps_vector.in
environment
tabular_data
  tabular_data_file = 'rosen_ps_vector.dat'

method
  vector_parameter_study
    final_point = 1.1 1.3
    num_steps = 10

variables
  continuous_design = 2
  initial_point = -0.3 0.2
  descriptors 'x1' 'x2'

interface
  analysis_driver = 'rosenbrock'
  direct

responses
  objective_functions = 1
  no_gradients
  no_hessians
```

Sample 5: Hybrid Strategy

The following sample input file shows a hybrid environment using three methods. It employs a genetic algorithm, pattern search, and full Newton gradient-based optimization in succession to solve the Textbook Example (see Textbook). A similar file is available as dakota/share/dakota/examples/users/textbook_hybrid_strat.in.

```dakota
environment
  hybrid sequential
    method_list = 'PS' 'PS2' 'NLP'

method
  id_method = 'PS'
  model_pointer = 'M1'
  coliny_pattern_search stochastic
    seed = 1234
    initial_delta = 0.1
    variable_tolerance = 1.e-4
    solution_accuracy = 1.e-10
    exploratory_moves basic_pattern
    #verbose output

method
  id_method = 'PS2'
  model_pointer = 'M1'
  max_function_evaluations = 10
  coliny_pattern_search stochastic
    seed = 1234
    initial_delta = 0.1
    variable_tolerance = 1.e-4
    solution_accuracy = 1.e-10
    exploratory_moves basic_pattern
    #verbose output

method
  id_method = 'NLP'
  model_pointer = 'M2'
```
optpp_newton
gradient_tolerance = 1.e-12
convergence_tolerance = 1.e-15
#verbose output

model
id_model = 'M1'
single
  variables_pointer = 'V1'
  interface_pointer = 'I1'
  responses_pointer = 'R1'
model
id_model = 'M2'
single
  variables_pointer = 'V1'
  interface_pointer = 'I1'
  responses_pointer = 'R2'

variables
id_variables = 'V1'
  continuous_design = 2
  initial_point = 0.6 0.7
  upper_bounds = 5.8 2.9
  lower_bounds = 0.5 -2.9
  descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
direct
  analysis_driver = 'text_book'

responses
id_responses = 'R1'
  objective_functions = 1
  no_gradients
  no_hessians
responses
id_responses = 'R2'
  objective_functions = 1
  analytic_gradients
  analytic_hessians

Additional example input files, as well as the corresponding output, are provided in the Tutorial chapter of the Users Manual [5].

5.4 Input Spec Summary

This file is derived automatically from dakota.xml, which is used in the generation of parser system files that are compiled into the Dakota executable. Therefore, these files are the definitive source for input syntax, capability options, and associated data inputs. Refer to the Developers Manual information on how to modify the input specification and propagate the changes through the parsing system.

Key features of the input specification and the associated user input files include:

- In the input specification, required individual specifications simply appear, optional individual and group specifications are enclosed in [], required group specifications are enclosed in (), and either-or relationships are denoted by the | symbol. These symbols only appear in dakota.input.summary; they must not appear in actual user input files.

- Keyword specifications (i.e., environment, method, model, variables, interface, and responses) begin with the keyword possibly preceded by white space (blanks, tabs, and newlines) both in the input specifications and in user input files. For readability, keyword specifications may be spread across several lines. Earlier versions of Dakota (prior to 4.1) required a backslash character (\) at the ends of intermediate lines of a keyword. While such backslashes are still accepted, they are no longer required.
• Some of the keyword components within the input specification indicate that the user must supply INTEGER, REAL, STRING, INTEGERLIST, REALLIST, or STRINGLIST data as part of the specification. In a user input file, the "=" is optional, data in a LIST can be separated by commas or whitespace, and the STRING data are enclosed in single or double quotes (e.g., 'text_book' or "text_book").

• In user input files, input is largely order-independent (except for entries in lists of data), case insensitive, and white-space insensitive. Although the order of input shown in the Sample Input Files generally follows the order of options in the input specification, this is not required.

• In user input files, specifications may be abbreviated so long as the abbreviation is unique. For example, the npsol_sqp specification within the method keyword could be abbreviated as npsol, but dot_sqp should not be abbreviated as dot since this would be ambiguous with other DOT method specifications.

• In both the input specification and user input files, comments are preceded by #.

• ALIAS refers to synonymous keywords, which often exist for backwards compatibility. Users are encouraged to use the most current keyword.

dakota.input.summary:

KEYWORD01 environment
  [ tabular_data ALIAS tabular_graphics_data
    [ tabular_data_file ALIAS tabular_graphics_file STRING ]
    [ custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    ]
    [ annotated ]
    [ freeform ]
  ]
  [ output_file STRING ]
  [ error_file STRING ]
  [ read_restart STRING
    [ stop_restart INTEGER >= 0 ]
  ]
  [ write_restart STRING ]
  [ output_precision INTEGER >= 0 ]
  [ results_output
    [ results_output_file STRING ]
    [ text ]
    [ hdf5 ]
    [ model_selection
      top_method
      | none
      | all_methods
      | all
    ]
    [ interface_selection
      none
      | simulation
      | all
    ]
  ]
  [ graphics ]
  [ check ]
  [ pre_run
    [ input STRING ]
    [ output STRING
      [ ( custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
      ]
      [ annotated ]
      [ freeform ]
    ]
  ]
| run |
| [ input STRING ] |
| [ output STRING ] |
| ] |
| post_run |
| [ input STRING |
| [ ( custom_annotated |
| [ header ] |
| [ eval_id ] |
| [ interface_id ] ] |
| | annotated |
| | freeform ] |
| ] |
| [ output STRING ] |
| ] |
| [ top_method_pointer ALIAS method_pointer STRING ] |

**KEYWORD method**

| id_method STRING |
| [ output debug |
| | verbose |
| | normal |
| | quiet |
| | silent ] |
| [ final_solutions INTEGER >= 0 ] |

| hybrid |
| ( sequential ALIAS uncoupled |
| ( method_name_list STRINGLIST |
| [ model_pointer_list STRINGLIST ] ] |
| | method_pointer_list STRINGLIST |
| [ iterator_servers INTEGER > 0 ] |
| [ iterator_scheduling |
| master |
| peer ] |
| [ processors_per_iterator INTEGER > 0 ] ] |
| | embedded ALIAS coupled |
| ( global_method_name STRING |
| [ global_model_pointer STRING ] ] |
| | [ global_method_pointer STRING ] |
| [ local_method_name STRING |
| [ local_model_pointer STRING ] ] |
| | [ local_method_pointer STRING |
| [ local_search_probability REAL ] |
| [ iterator_servers INTEGER > 0 ] |
| [ iterator_scheduling |
| master |
| peer ] |
| [ processors_per_iterator INTEGER > 0 ] ] |
| | collaborative |
| ( method_name_list STRINGLIST |
| [ model_pointer_list STRINGLIST ] ] |
| | [ method_pointer_list STRINGLIST |
| [ iterator_servers INTEGER > 0 ] |
| [ iterator_scheduling |
| master |
| peer ] |
| [ processors_per_iterator INTEGER > 0 ] |
5.4 Input Spec Summary

```dakota

multi_start
  method_name STRING
  method_pointer STRING
  random_starts INTEGER
  seed INTEGER
  starting_points REALLIST
  iterator_servers INTEGER > 0
  iterator_scheduling
    master
    peer
  processors_per_iterator INTEGER > 0

pareto_set
  method_name ALIAS opt_method_name STRING
  method_pointer ALIAS opt_method_pointer STRING
  random_weight_sets INTEGER
  seed INTEGER
  weight_sets ALIAS multi_objective_weight_sets REALLIST
  iterator_servers INTEGER > 0
  iterator_scheduling
    master
    peer
  processors_per_iterator INTEGER > 0

branch_and_bound
  method_pointer STRING
  method_name STRING
  scaling

surrogate_based_local
  method_pointer ALIAS approx_method_pointer STRING
  method_name ALIAS approx_method_name STRING
  model_pointer ALIAS approx_model_pointer STRING
  soft_convergence_limit INTEGER
  truth_surrogate_bypass
  approx_subproblem
    original_primary
    single_objective
    augmented_lagrangian_objective
    lagrangian_objective
    original_constraints
    linearized_constraints
    no_constraints
  merit_function
    penaltyMerit
    adaptive_penalty_merit
    lagrangianMerit
    augmented_lagrangian_merit
  acceptance_logic
    tr_ratio
    filter
  constraint_relax
  homotopy
```
[ trust_region
  [ initial_size REALIST ]
  [ minimum_size REAL ]
  [ contract_threshold REAL ]
  [ expand_threshold REAL ]
  [ contraction_factor REAL ]
  [ expansion_factor REAL ]
]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
)
)
| surrogate_based_global
  method_pointer ALIAS approx_method_pointer STRING
  | method_name ALIAS approx_method_name STRING
  model_pointer ALIAS approx_model_pointer STRING
  [ replace_points ]
  [ max_iterations INTEGER >= 0 ]
)
| dot_frcg
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)
| dot_mmfd
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)
| dot_bfgs
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)
| dot_slp
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)
| dot_sqp
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)
| conmin_frcg
5.4 Input Spec Summary

[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| (conmin_mfd
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| (dl_solver STRING
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| (npsol_sqp
 [ verify_level INTEGER ]
 [ function_precision REAL ]
 [ linesearch_tolerance REAL ]
 [ convergence_tolerance REAL ]
 [ max_iterations INTEGER >= 0 ]
 [ constraint_tolerance REAL ]
 [ speculative ]
 [ max_function_evaluations INTEGER >= 0 ]
 [ scaling ]
 [ model_pointer STRING ]
)

| (nlssol_sqp
 [ verify_level INTEGER ]
 [ function_precision REAL ]
 [ linesearch_tolerance REAL ]
 [ convergence_tolerance REAL ]
 [ max_iterations INTEGER >= 0 ]
 [ constraint_tolerance REAL ]
 [ speculative ]
 [ max_function_evaluations INTEGER >= 0 ]
 [ scaling ]
 [ model_pointer STRING ]
)

| (nlpq1_sqp
 [ max_iterations INTEGER >= 0 ]
 [ convergence_tolerance REAL ]
 [ max_function_evaluations INTEGER >= 0 ]
 [ scaling ]
 [ model_pointer STRING ]
)

| (optpp_cg
 [ max_step REAL ]
 [ gradient_tolerance REAL ]
 [ max_iterations INTEGER >= 0 ]
 [ convergence_tolerance REAL ]
 [ speculative ]
 [ max_function_evaluations INTEGER >= 0 ]
 [ scaling ]
 [ model_pointer STRING ]
)

| (optpp_q_newton
 [ search_method
 value_based_line_search
]
gradient_based_line_search
| trust_region
| tr_pds
|
merit_function
| el_bakry
| argaez_tapia
| van_shanno
|
| steplength_to_boundary REAL |
| centering_parameter REAL |
| max_step REAL |
| gradient_tolerance REAL |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| speculative |
| max_function_evaluations INTEGER >= 0 |
| scaling |
| model_pointer STRING |

|
| optpp_fd_newton
| search_method
| value_based_line_search
| gradient_based_line_search
| trust_region
| tr_pds
|
| merit_function
| el_bakry
| argaez_tapia
| van_shanno
|
| steplength_to_boundary REAL |
| centering_parameter REAL |
| max_step REAL |
| gradient_tolerance REAL |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| speculative |
| max_function_evaluations INTEGER >= 0 |
| scaling |
| model_pointer STRING |

|
| optpp_g_newton
| search_method
| value_based_line_search
| gradient_based_line_search
| trust_region
| tr_pds
|
| merit_function
| el_bakry
| argaez_tapia
| van_shanno
|
| steplength_to_boundary REAL |
| centering_parameter REAL |
| max_step REAL |
| gradient_tolerance REAL |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| speculative |
| max_function_evaluations INTEGER >= 0 |
| scaling |
| model_pointer STRING |

|
| optpp_newton
| search_method
| value_based_line_search
| gradient_based_line_search
| trust_region
| tr_pds
|
| trust_region
| tr_pds |

[ merit_function
  el_bakry
  | argaez_tapia
  | van_shanno |

[ steplength_to_boundary REAL ]
[ centering_parameter REAL ]
[ max_step REAL ]
[ gradient_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

| optpp_pds

[ search_scheme_size INTEGER ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

| demo_tpl

[ max_function_evaluations INTEGER >= 0 ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ variable_tolerance REAL ]
[ solution_target ALIAS solution_accuracy REAL ]
[ options_file STRING ]

| rol

[ max_iterations INTEGER >= 0 ]
[ variable_tolerance REAL ]
[ gradient_tolerance REAL ]
[ constraint_tolerance REAL ]
[ options_file STRING ]
[ scaling ]
[ model_pointer STRING ]

| asynch_pattern_search ALIAS coliny_apps

[ initial_delta REAL ]
[ contraction_factor REAL ]
[ variable_tolerance REAL ]
[ solution_target ALIAS solution_accuracy REAL ]
[ synchronization
  blocking
  | nonblocking |

[ merit_function
  merit_max
  | merit_max_smooth
  | merit1
  | merit1_smooth
  | merit2
  | merit2_smooth
  | merit2_squared |

[ constraint_penalty REAL ]
[ smoothing_factor REAL ]
[ constraint_tolerance REAL ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

<p>|</p>
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<td>function_precision REAL</td>
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<td>display_format STRING</td>
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</tr>
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<tr>
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<td>scaling</td>
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<td>model_pointer STRING</td>
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</tr>
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<td>expand_threshold REAL</td>
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<tr>
<td>contraction_factor REAL</td>
</tr>
<tr>
<td>expansion_factor REAL</td>
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<tr>
<td>max_iterations INTEGER &gt;= 0</td>
</tr>
<tr>
<td>max_function_evaluations INTEGER &gt;= 0</td>
</tr>
<tr>
<td>scaling</td>
</tr>
<tr>
<td>model_pointer STRING</td>
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</tbody>
</table>
|}

<table>
<thead>
<tr>
<th>snowpac</th>
</tr>
</thead>
<tbody>
<tr>
<td>seed INTEGER &gt; 0</td>
</tr>
<tr>
<td>trust_region</td>
</tr>
<tr>
<td>initial_size REALLIST</td>
</tr>
<tr>
<td>minimum_size REAL</td>
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<tr>
<td>contract_threshold REAL</td>
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<tr>
<td>contraction_factor REAL</td>
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<td>expansion_factor REAL</td>
</tr>
<tr>
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<tr>
<td>max_function_evaluations INTEGER &gt;= 0</td>
</tr>
<tr>
<td>scaling</td>
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<td>model_pointer STRING</td>
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<td>( below_limit REAL</td>
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<td>distance REALLIST</td>
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5.4 Input Spec Summary

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  [ percent_change REAL ]
  [ num_generations INTEGER >= 0 ]
] [ postprocessor_type
  orthogonl_distance REALIST
  ] [ max_iterations INTEGER >= 0 ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ population_size INTEGER >= 0 ]
  [ log_file STRING ]
  [ print_each_pop ]
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    | unique_random
    | flat_file STRING
  ] [ crossover_type
    multi_point_binary INTEGER
    | multi_point_parameterized_binary INTEGER
    | multi_point_real INTEGER
    | ( shuffle_random
      [ num_parents INTEGER > 0 ]
      [ num_offspring INTEGER > 0 ]
    )
    [ crossover_rate REAL ]
  ] [ mutation_type
    bit_random
    | replace_uniform
    |
    ( offset_normal
      [ mutation_scale REAL ]
    )
    |
    ( offset_cauchy
      [ mutation_scale REAL ]
    )
    |
    ( offset_uniform
      [ mutation_scale REAL ]
    )
    [ mutation_rate REAL ]
  ] [ seed INTEGER > 0 ]
  [ convergence_tolerance REAL ]
  [ model_pointer STRING ]
] [ soga
  [ fitness_type
    merit_function
    [ constraint_penalty REAL ]
  ] [ replacement_type
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    | favor_feasible
    | roulette_wheel
    | unique_roulette_wheel
  ] [ convergence_type
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      [ percent_change REAL ]
      [ num_generations INTEGER >= 0 ]
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    |
    ( average_fitness_tracker
      [ percent_change REAL ]
      [ num_generations INTEGER >= 0 ]
    )
  ]
}
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ population_size INTEGER >= 0 ]
[ log_file STRING ]
[ print_each_pop ]
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  | unique_random
  | flat_file STRING ]
[ crossover_type
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  | multi_point_parameterized_binary INTEGER
  | multi_point_real INTEGER
  | shuffle_random
  [ num_parents INTEGER > 0 ]
  [ num_offspring INTEGER > 0 ]
]  
[ crossover_rate REAL ]
]
[ mutation_type
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  | replace_uniform
  | ( offset_normal
    [ mutation_scale REAL ]
  )
  | ( offset_cauchy
    [ mutation_scale REAL ]
  )
  | ( offset_uniform
    [ mutation_scale REAL ]
  )
  [ mutation_rate REAL ]
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[ seed INTEGER > 0 ]
[ convergence_tolerance REAL ]
[ model_pointer STRING ]
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[ coliny_pattern_search
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    | simplex
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  [ stochastic ]
  [ total_pattern_size INTEGER ]
[ exploratory_moves
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  | adaptive_pattern
  | basic_pattern
]  
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  | nonblocking
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[ constraint_penalty REAL ]
[ initial_delta REAL ]
[ variable_tolerance REAL ]
[ solution_target ALIAS solution_accuracy REAL ]
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[ misc_options STRINGLIST ]
5.4 Input Spec Summary

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[ coliny_direct
[ division
  major_dimension
  | all_dimensions
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[ local_balance_parameter REAL ]
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[ min_boxsize_limit REAL ]
[ constraint_penalty REAL ]
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[ convergence_tolerance REAL ]
[ max_function_evaluations INTEGER >= 0 ]
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[ model_pointer STRING ]
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  | unique_random
  | flat_file STRING
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[ fitness_type
  linear_rank
  | merit_function
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Dakota Input Specification

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  | chc INTEGER
  | elitist INTEGER
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  | blend
  | uniform
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    [ mutation_range INTEGER ]
  )
  |
  ( offset_cauchy
    [ mutation_scale REAL ]
    [ mutation_range INTEGER ]
  )
  |
  ( offset_uniform
    [ mutation_scale REAL ]
    [ mutation_range INTEGER ]
  )
  [ non_adaptive ]
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[ constraint_penalty REAL ]
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  [ convergence_tolerance REAL ]
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| ( nl2sol
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  [ absolute_conv_tol REAL ]
  [ x_conv_tol REAL ]
  [ singular_conv_tol REAL ]
  [ singular_radius REAL ]
  [ false_conv_tol REAL ]
  [ initial_trust_radius REAL ]
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5.4 Input Spec Summary

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    [exploration INTEGER >= 0]  
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    [blocking]  
    [nonblocking]
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  [x_conv_tol REAL]  
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    (surfpack  
      [export_model]  
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        [text_archive]  
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    )
  ]
)

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Dakota Input Specification

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| freeform |
| active_only |
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[ model_pointer STRING ]

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| random |
[ rng
mt19937
| rnum2 |
[ probability_refinement ALIAS sample_refinement
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| adapt_import
| mm_adapt_import
[ refinement_samples INTEGERLIST ]
| ]
[ final_moments
none
| standard
| central |
| ]

[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
probabilities
| reliabilities
| gen_reliabilities
[ system
series
| parallel
] ]

[ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
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[ reliability_levels REALLIST
[ num_reliability_levels INTEGERLIST ]
]

[ gen_reliability_levels REALLIST
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[ distribution
cumulative
| complementary |
| variance_based_decomp
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      | increment_start_order
      | increment_max_rank
      | increment_max_order
      | increment_max_rank_order
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    | absolute
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  [ tensor_grid ]
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  [ arithmetic_tolerance REAL ]
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  [ max_order INTEGER >= 0 ]
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    | gen_reliabilities
    | system
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[ distribution
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  | complementary
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| increment_start_order
| increment_max_rank
| increment_max_order
| increment_max_rank_order
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| convergence_tolerance REAL ]
| metric_scale
| relative
| absolute
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| statistics_mode
| active
| combined
|
| allocation_control
| greedy
|
| discrepancy_emulation
| distinct ALIAS paired
| recursive
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| rounding_tolerance REAL ]
| arithmetic_tolerance REAL ]
| regression_type
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| max_cross_iterations INTEGER >= 0 ]
| solver_tolerance REAL ]
| response_scaling ]
| tensor_grid ]
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| collocation_ratio REAL ]
| start_order_sequence ALIAS order_sequence INTEGERLIST ]
| dimension_preference REALLIST ]
| ]
| adapt_order ]
| kick_order INTEGER > 0 ]
| max_order INTEGER >= 0 ]
| max_cv_order_candidates INTEGER >= 0 ]
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| kick_rank INTEGER > 0 ]
| max_rank INTEGER >= 0 ]
| max_cv_rank_candidates INTEGER >= 0 ]
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| random
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| rng
| mt19937
| rnum2
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| adapt_import
| mm_adapt_import
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| ]
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| standard
| central
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[ compute
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  | reliabilities
  | gen_reliabilities
  [ system
  series
  | parallel
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[ probability_levels REALLIST
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[ reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ]
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[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
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[ distribution
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  | complementary
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    [ interface_id ]
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[ model_pointer STRING ]
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[ arithmetic_tolerance REAL ]
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[ max_order INTEGER >= 0 ]
[ max_cv_order_candidates INTEGER >= 0 ]
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[ max_rank INTEGER >= 0 ]
[ max_cv_rank_candidates INTEGER >= 0 ]
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| random
| ]
[ rng
| mt19937
| rnum2
| ]
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| mm_adapt_import
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| final_moments
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| standard
| central
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| [ num_response_levels INTEGERLIST ]
| [ compute
| probabilities
| | reliabilities
| | gen_reliabilities
| | system
| | series
| | parallel
| ]
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| [ num_probability_levels INTEGERLIST ]
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| reliability_levels REALLIST
| [ num_reliability_levels INTEGERLIST ]
|
| gen_reliability_levels REALLIST
| [ num_gen_reliability_levels INTEGERLIST ]
|
| distribution
| cumulative
| | complementary
|
| variance_based_decomp
| [ interaction_order INTEGER > 0 ]
[ drop_tolerance REAL ]
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| full_covariance ]
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| eval_id ]
| interface_id ]
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| annotated
| freeform ]
[ active_only ]
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| eval_id ]
| interface_id ]
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| freeform ]
]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]
]

[ polynomial_chaos ALIAS nond_polynomial_chaos
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| ( dimension_adaptive
| sobol
| decay
| generalized
)
]
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[ convergence_tolerance REAL ]
[ metric_scale
| relative
| absolute
]
[ quadrature_order INTEGER
[ dimension_preference REALLIST ]
[ nested ]
| non_nested ]
]

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[ dimension_preference REALLIST ]
[ restricted ]
| unrestricted ]
| nested
| non_nested ]
]
[ cubature_integrand INTEGER
]

[ expansion_order INTEGER
[ dimension_preference REALLIST ]
[ basis_type
| tensor_product
| total_order
| ]
| ( adapted
[ advancements INTEGER ]
[ soft_convergence_limit INTEGER ]
)
]
[ collocation_points INTEGER
[ ( least_squares
| svd
]
5.4 Input Spec Summary

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[ interface_id ]
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| annotated
| freeform
[ active_only ]
)
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{ orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
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| tensor_grid INTEGERLIST ]
| reuse_points ALIAS reuse_samples ]
| import_build_points_file ALIAS import_points_file STRING
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| eval_id ]
| interface_id ]
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| annotated
| freeform
| active_only ]
)
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| wiener ]
| normalized ]
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| rng
| mt19937
| rnum2 ]
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| adapt_import
| mm_adapt_import
| refinement_samples INTEGERLIST ]
|
| final_moments
| none
| standard
| central
]
| response_levels REALLIST
| num_response_levels INTEGERLIST ]
| compute
| probabilities
| reliabilities
| gen_reliabilities
| system
| series
| parallel
|
|
| probability_levels REALLIST
| num_probability_levels INTEGERLIST ]
|
| reliability_levels REALLIST
| num_reliability_levels INTEGERLIST ]
|
| gen_reliability_levels REALLIST
| num_gen_reliability_levels INTEGERLIST ]
|
| distribution
| cumulative
| complementary
5.4 Input Spec Summary

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  [ drop_tolerance REAL ]
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  | full_covariance ]
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  | freeform ]
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[ fixed_seed ]
[ model_pointer STRING ]
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      |
      ( dimension_adaptive
        sobol
        | decay
        | generalized
      )
    ]
  ]
[ max_refinement_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ metric_scale
  relative
  | absolute
]
[ statistics_mode
  active
  | combined
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[ allocation_control
  greedy
]
[ discrepancy_emulation
  distinct ALIAS paired
  | recursive
]
[ quadrature_order_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  [ nested
    | non_nested ]
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  [ dimension_preference REALLIST ]
  [ restricted
    | unrestricted ]
  [ nested
    | non_nested ]
]
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[ dimension_preference REALLIST ]
[ basis_type
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  | total_order
  | ( adapted
  [ advancements INTEGER ]
  [ soft_convergence_limit INTEGER ]
 ]
[ collocation_ratio REAL
  [ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
  [ least_squares
    | svd
    | equality_constrained ]
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[ orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
  | basis_pursuit ALIAS bp
  | ( basis_pursuit_denoising ALIAS bpdn
  [ noise_tolerance REALLIST ]
 ]
[ least_angle_regression ALIAS lars
  [ noise_tolerance REALLIST ]
 ]
[ least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]
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[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
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 ]
[ import_build_points_file ALIAS import_points_file STRING
  [ custom_annotated
    header
    | eval_id
    | interface_id
  ]
  | annotated
  | freeform
  | active_only
 ]
]
[ orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
  [ import_build_points_file ALIAS import_points_file STRING
    [ custom_annotated
      header
      | eval_id
      | interface_id
    ]
    | annotated
    | freeform
  ]
]
5.4 Input Spec Summary

[ active_only ]

[ askey
| wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type
  lhs
  | random ]
[ rng
  mt19937
  | rnum2 ]
[ probability_refinement ALIAS sample_refinement
  import
  | adapt_import
  | mm_adapt_import
  [ refinement_samples INTEGERLIST ] ]
[ final_moments
  none
  | standard ]
| central ]
[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | reliabilities
    | gen_reliabilities
    [ system
      series
      | parallel ] ]
]
[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ] ]
[ reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ] ]
[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ] ]
[ distribution
  cumulative ]
| complementary ]
[ variance_based_decomp
  [ interaction_order INTEGER > 0 ]
  [ drop_tolerance REAL ] ]
[ diagonal_covariance
  | full_covariance ]
[ import_approx_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ] ]
[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ] ]
| annotated |
| freeform |

| seed_sequence INTEGERLIST |
| fixed_seed |
| model_pointer STRING |

| multilevel_polynomial_chaos |
| max_iterations INTEGER >= 0 |
| allocation_control |
| estimator_variance |
| estimator_rate REAL |
| rip_sampling |
| convergence_tolerance REAL |
| metric_scale |
| relative |
| absolute |
| discrepancy_emulation |
| distinct ALIAS paired |
| recursive |

| expansion_order_sequence INTEGERLIST |
| dimension_preference REALLIST |
| basis_type |
| tensor_product |
| total_order |
| (adapted |
| advancements INTEGER |
| soft_convergence_limit INTEGER |
| ) |

| collocation_ratio REAL |
| collocation_points_sequence ALIAS pilot_samples INTEGERLIST |
| (least_squares |
| svd |
| equality_constrained |
| ) |
| (orthogonal_matching_pursuit ALIAS omp |
| noise_tolerance REALLIST |
| ) |
| basis_pursuit ALIAS bp |
| (basis_pursuit_denoising ALIAS bpdn |
| noise_tolerance REALLIST |
| ) |
| (least_angle_regression ALIAS lars |
| noise_tolerance REALLIST |
| ) |
| (least_absolute_shrinkage ALIAS lasso |
| noise_tolerance REALLIST |
| l2_penalty REAL |
| ) |
| cross_validation |
| noise_only |
| max_cv_order_candidates INTEGER >= 0 |

| ratio_order REAL |
| response_scaling |
| use_derivatives |
| tensor_grid |
| reuse_points ALIAS reuse_samples |
| max_solver_iterations INTEGER >= 0 |


{ expansion_samples_sequence INTEGERLIST
   [ reuse_points ALIAS reuse_samples ]
 }

[ import_build_points_file ALIAS import_points_file STRING
   [{ custom_annotated
     [ header ]
     [ eval_id ]
     [ interface_id ]
   }]
   | annotated
   | freeform
   [ active_only ]
]

{ orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
collocation_points_sequence INTEGERLIST
   [ tensor_grid INTEGERLIST ]
   [ reuse_points ALIAS reuse_samples ]
   [ import_build_points_file ALIAS import_points_file STRING
     [{ custom_annotated
       [ header ]
       [ eval_id ]
       [ interface_id ]
     }]
     | annotated
     | freeform
     [ active_only ]
   ]
   | askey
   | wiener
   | normalized
   [ export_expansion_file STRING ]
   [ samples_on_emulator ALIAS samples INTEGER ]
   [ sample_type
     lhs
     | random
   ]
   [ rng
     mt19937
     | rnum2
   ]
   [ probability_refinement ALIAS sample_refinement
     import
     | adapt_import
     | mm_adapt_import
     [ refinement_samples INTEGERLIST ]
   ]
   [ final_moments
     none
     | standard
     | central
   ]
   [ response_levels REALLIST
     [ num_response_levels INTEGERLIST ]
     [ compute
       probabilities
       | reliabilities
       | gen_reliabilities
       [ system
         series
         | parallel
       ]
     ]
   ]
   [ probability_levels REALLIST
     [ num_probability_levels INTEGERLIST ]
   ]
   [ reliability_levels REALLIST
     [ num_reliability_levels INTEGERLIST ]
   ]
   [ gen_reliability_levels REALLIST
   ]
[ num_gen_reliability_levels INTEGERLIST ]
]

distribution
cumulative
| complementary
]
]
[ variance_based_decomp
[ interaction_order INTEGER > 0 ]
[ drop_tolerance REAL ]
]
]
diagonal_covariance
| full_covariance
]
[ import_approx_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform
| active_only
]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform
]
]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]
]
]
[ stoch_collocation ALIAS nond_stoch_collocation
[ ( p_refinement
  uniform
  | ( dimension_adaptive
    sobol
    | generalized
  )
  )
]
[ h_refinement
  uniform
  | ( dimension_adaptive
    sobol
    | generalized
  )
  | local_adaptive
  ]
[ max_refinement_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ metric_scale
  relative
  | absolute
  ]
[ quadrature_order INTEGER
  [ dimension_preference REALLIST ]
  [ nested
  | non_nested ]
  ]
]
[ sparse_grid_level INTEGER
  [ dimension_preference REALLIST ]
  [ nodal
  | hierarchical
  | restricted
  | unrestricted ]
]
5.4 Input Spec Summary

```plaintext
[ nested
 | non_nested ]
)
[ piecewise
 | askey
 | wiener ]
[ use_derivatives ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type
 lhs
 | random
 ]
[ rng
 mt19937
 | rnum2
]
[ probability_refinement ALIAS sample_refinement
 import
 | adapt_import
 | mm_adapt_import
 [ refinement_samples INTEGERLIST ]
]
[ final_moments
 none
 | standard
 | central
]
[ response_levels REALLIST
 [ num_response_levels INTEGERLIST ]
 [ compute
 probabilities
 | reliabilities
 | gen_reliabilities
 [ system
 series
 | parallel
 ]
 ]
]
[ probability_levels REALLIST
 [ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
 [ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
 [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
 cumulative
 | complementary
]
[ variance_based_decomp
 [ interaction_order INTEGER > 0 ]
 [ drop_tolerance REAL ]
]
[ diagonal_covariance
 | full_covariance ]
[ import_approx_points_file STRING
 [ ( custom_annotated
 [ header ]
 [ eval_id ]
 [ interface_id ]
 )
 | annotated
 | freeform ]
 | active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
 [ ( custom_annotated
 [ header ]
 [ eval_id ]
 [ interface_id ]
 )
]```
multifidelity_stoch_collocation
|
| p_refinement
| uniform
| ( dimension_adaptive
| sobol
| generalized
|
| h_refinement
| uniform
| ( dimension_adaptive
| sobol
| generalized
| local_adaptive
|
| max_refinement_iterations INTEGER >= 0
| convergence_tolerance REAL
| metric_scale
| relative
| absolute
|
| statistics_mode
| active
| combined
|
| allocation_control
| greedy
|
| discrepancy_emulation
| distinct ALIAS paired
| recursive
|
| quadrature_order_sequence INTEGERLIST
| dimension_preference REALLIST
| nested
| non_nested
|
| sparse_grid_level_sequence INTEGERLIST
| dimension_preference REALLIST
| nodal
| hierarchical
| restricted
| unrestricted
| nested
| non_nested
|
| piecewise
| askey
| wiener
| use_derivatives
| samples_on_emulator ALIAS samples INTEGER
| sample_type
| lbs
| random
|
| rng
| mt19937
| rnum2
[ probability_refinement ALIAS sample_refinement
    import
    | adapt_import
    | mm_adapt_import
    [ refinement_samples INTEGERLIST ]
]
[ final_moments
    none
    | standard
    | central
]
[ response_levels REALLIST
    [ num_response_levels INTEGERLIST ]
    [ compute
        probabilities
        | reliabilities
        | gen_reliabilities
        [ system
            series
            | parallel
        ]
    ]
][ probability_levels REALLIST
    [ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
    [ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
    [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
    cumulative
    | complementary
]
[ variance_based_decomp
    [ interaction_order INTEGER > 0 ]
    [ drop_tolerance REAL ]
]
[ diagonal_covariance
    | full_covariance
][ import_approx_points_file STRING
    [ ( custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
    )
    | annotated
    | freeform
    | active_only
]
[ export_approx_points_file ALIAS export_points_file STRING
    [ ( custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
    )
    | annotated
    | freeform
]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]
]
[ sampling ALIAS nond_sampling
    [ samples ALIAS initial_samples INTEGER ]
    [ seed INTEGER > 0 ]
    [ fixed_seed ]
    [ sample_type
        lhs
        | random
]
incremental_lhs
incremental_random

refinement_samples INTEGERLIST

d_optimal
candidate_designs INTEGER > 0
leja_oversample_ratio REAL

variance_based_decomp
drop_tolerance REAL

backfill

principal_components
percent_variance_explained REAL

wilks
order INTEGER
candidate_level REAL
one_sided_lower
one_sided_upper
two_sided

final_moments
none
standard
central

response_levels REALLIST
num_response_levels INTEGERLIST
compute
probabilities
reliabilities
gen_reliabilities
system
series
parallel

probability_levels REALLIST
num_probability_levels INTEGERLIST

reliability_levels REALLIST
num_reliability_levels INTEGERLIST

gen_reliability_levels REALLIST
num_gen_reliability_levels INTEGERLIST

distribution
cumulative
complementary

rng
mt19937
rnum2

model_pointer STRING

multilevel_sampling ALIAS multilevel_mc ALIAS mlmc
seed_sequence INTEGERLIST
fixed_seed
pilot_samples ALIAS initial_samples INTEGERLIST
sample_type
lhs
random

export_sample_sequence
(custom_annotated
header
eval_id
interface_id)
5.4 Input Spec Summary

| annotated |
| freeform |

| allocation_target |
| mean |
| variance |
| standard_deviation |
| scalarization |

| qoi_aggregation |
| sum |
| max |

| convergence_tolerance |
| convergence_tolerance_type |
| relative |
| absolute |

| convergence_tolerance_target |
| variance_constraint |
| cost_constraint |

| max_iterations |
| max_function_evaluations |
| final_moments |
| none |
| standard |
| central |

| distribution |
| cumulative |
| complementary |

| rng |
| mt19937 |
| rnum2 |

| model_pointer |
| STRING |

| multifidelity_sampling |
| ALIAS multifidelity_mc |
| ALIAS mfmc |

| seed_sequence |
| fixed_seed |

| pilot_samples |
| ALIAS initial_samples |

| solution_mode |
| online_pilot |
| offline_pilot |
| pilot_projection |

| sample_type |
| lhs |
| random |

| export_sample_sequence |
| ( custom.annotated |
| header |
| eval_id |
| interface_id |
| annotated |
| freeform |

| convergence_tolerance |
| REAL |
Dakota Input Specification

[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ final_moments
  none
  | standard
  | central
 ]
[ distribution
  cumulative
  | complementary
 ]
[ rng
  mt19937
  | rnum2
 ]
[ model_pointer STRING ]
)

| multilevel_multifidelity_sampling ALIAS multilevel_multifidelity_mc ALIAS mlmfmc
  [ seed_sequence INTEGERLIST ]
  [ fixed_seed ]
  [ pilot_samples ALIAS initial_samples INTEGERLIST ]
  [ sample_type
    lhs
    | random
  ]
  [ export_sample_sequence
    [ ( custom.annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
    | annotated
    | freeform
  ]
  [ convergence_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ final_moments
    none
    | standard
    | central
  ]
  [ distribution
    cumulative
    | complementary
  ]
  [ rng
    mt19937
    | rnum2
  ]
  [ model_pointer STRING ]
)

| approximate_control_variate ALIAS acv_sampling
  acv.independent_sampling ALIAS acv_is
  | acv_multifidelity ALIAS acv_mf
  | acv_kl ALIAS acv_adaptive
  [ pilot_samples ALIAS initial_samples INTEGERLIST ]
  [ solution_mode
    online_pilot
    | offline_pilot
    | pilot_projection
  ]
  [ truth_fixed_by_pilot ]
  [ sqp
    | nip ]
  [ seed_sequence INTEGERLIST ]
  [ fixed_seed ]
  [ sample_type
    lhs
    | random
  ]

[ export_sample_sequence
  [ [ custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  ]
  ]
  [ annotated
  [ freeform ]
]
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ final_moments
  none
  | standard
  | central
]
[ distribution
  cumulative
  | complementary
]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)
)

[ importance_sampling ALIAS nond_importance_sampling
[ samples ALIAS initial_samples INTEGER ]
[ seed INTEGER > 0 ]
import
  | adapt import
  | mm_adapt_import
[ refinement_samples INTEGERLIST ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  ]
[ compute
  probabilities
  | gen_reliabilities
  [ system
    series
    | parallel
  ]
]
[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  cumulative
  | complementary
]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)
)

[ gpais ALIAS gaussian_process_adaptive_importance_sampling
[ build_samples ALIAS samples INTEGER ]
[ seed INTEGER > 0 ]
[ samples_on_emulator INTEGER ]
[ import_build_points_file ALIAS import_points_file STRING
  [ [ custom_annotated
    [ header ]
    [ eval_id ]
  ]
  ]
]
Dakota Input Specification

Dakota Version 6.15 Reference Manual generated on November 5, 2021

```plaintext
[ interface_id ]
| annotated
| freeform |
[ active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
]
[ max_iterations INTEGER >= 0 ]
[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | gen_reliabilities
    | system
    series
    | parallel
  ]
]
[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  cumulative
  | complementary
]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)
( adaptive_sampling ALIAS nond_adaptive_sampling
[ initial_samples ALIAS samples INTEGER ]
[ seed INTEGER > 0 ]
[ samples_on_emulator INTEGER ]
[ fitness_metric
  predicted_variance
  | distance
  | gradient
]
[ batch_selection
  naive
  | distance_penalty
  | topology
  | constant_liar
]
[ refinement_samples INTEGERLIST ]
[ import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
    [ header ]
```
5.4 Input Spec Summary

[ eval_id ]
[ interface_id ]
| annotated
| freeform |
[ misc_options STRINGLIST ]
[ max_iterations INTEGER >= 0 ]
[ response_levels REALLIST ]
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| gen_reliabilities
[ system
series
| parallel
]
]
[ probability_levels REALLIST ]
[ num_probability_levels INTEGERLIST ]
[ gen_reliability_levels REALLIST ]
[ num_gen_reliability_levels INTEGERLIST ]
[ distribution
cumulative
| complementary
]
[ rng
mt19937
| rnum2
]
[ model_pointer STRING ]
)
|
[ pof_darts ALIAS nond_pof_darts
build_samples ALIAS samples INTEGER
| seed INTEGER > 0 ]
[ lipschitz
local
| global
]
[ samples_on_emulator INTEGER ]
[ response_levels REALLIST ]
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| gen_reliabilities
[ system
series
| parallel
]
]
[ probability_levels REALLIST ]
[ num_probability_levels INTEGERLIST ]
[ gen_reliability_levels REALLIST ]
[ num_gen_reliability_levels INTEGERLIST ]
[ distribution
cumulative
| complementary
]
[ rng
mt19937
| rnum2
]
[ model_pointer STRING ]
)
|
[ rkd_darts ALIAS nond_rkd_darts
build_samples ALIAS samples INTEGER
| seed INTEGER > 0 ]
[ lipschitz
local
| global
]
[ samples_on_emulator INTEGER ]
[ response_levels REALLIST ]
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| gen_reliabilities
[ system
series
| parallel
]
]
[ probability_levels REALLIST ]
[ num_probability_levels INTEGERLIST ]
[ gen_reliability_levels REALLIST ]
[ num_gen_reliability Levels INTEGERLIST ]
[ distribution
cumulative
| complementary
]
[ rng
mt19937
| rnum2
]
[ model_pointer STRING ]
)
|
{ rkd_darts ALIAS nond_rkd_darts
build_samples ALIAS samples INTEGER
   [ seed INTEGER > 0 ]
   [ lipschitz
     local
     | global
   ]
   [ samples_on_emulator INTEGER ]
response_levels REALLIST
   [ num_response_levels INTEGERLIST ]
   [ compute
     probabilities
     | gen_reliabilities
     [ system
       series
     | parallel
   ]
   ]
probability_levels REALLIST
   [ num_probability_levels INTEGERLIST ]
[ gen_reliability_levels REALLIST
   [ num_gen_reliability_levels INTEGERLIST ]
]distribution
cumulative
| complementary
]rng
t19937
| rnum2
[model_pointer STRING ]
)(
global_evidence ALIAS nond_global_evidence
   [ samples INTEGER ]
   [ seed INTEGER > 0 ]
  [ ( sbo
     [ gaussian_process ALIAS kriging
       surfpack
       | dakota
     ]
     [ use_derivatives ]
     [ import_build_points_file ALIAS import_points_file STRING
       [ ( custom.annotated
          [ header ]
          [ eval_id ]
          [ interface_id ]
        )
        | annotated
        | freeform
        [ active_only ]
      ]
      [ export_approx_points_file ALIAS export_points_file STRING
        [ ( custom.annotated
          [ header ]
          [ eval_id ]
          [ interface_id ]
        )
        | annotated
        | freeform
      ]
    ]
  ]
  [ ego
    [ gaussian_process ALIAS kriging
      surfpack
      | dakota
    ]
    [ use_derivatives ]
    [ import_build_points_file ALIAS import_points_file STRING
      [ ( custom.annotated
    ]
  ]
)
Dakota Version 6.15 Reference Manual generated on November 5, 2021

5.4 Input Spec Summary

```
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform
| active_only
]

| export_approx_points_file ALIAS export_points_file STRING
|
|
{| custom_annotated
| header |
| eval_id |
| interface_id |
)
| annotated
| freeform
|
]
| ea
| lhs |
| response_levels REALLIST
|
| num_response_levels INTEGERLIST |
| compute
| probabilities
| gen_reliabilities
| system
| series
| parallel
|
|
| probability_levels REALLIST
| num_probability_levels INTEGERLIST |
|
| gen_reliability_levels REALLIST
| num_gen_reliability_levels INTEGERLIST |
| distribution
| cumulative
| complementary
|
| rng
| mt19937
| rnum2
|
| model_pointer STRING |
)
|
| global_interval_est ALIAS nond_global_interval_est
|
| samples INTEGER |
| seed INTEGER > 0 |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| max_function_evaluations INTEGER >= 0 |
| sho
| gaussian_process ALIAS kriging
| surfpack
| dakota
|
| experimental
| options_file STRING |
|
| use_derivatives |
|
| import_build_points_file ALIAS import_points_file STRING
| custom_annotated
| header |
| eval_id |
| interface_id |
|
| annotated
| freeform
| active_only
```
Dakota Input Specification

[ export_approx_points_file ALIAS export_points_file STRING
  ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
| annotated
| freeform
]

| ego
[ gaussian_process ALIAS kriging
  ( surfpack
    [ export_model
      [ filename_prefix STRING ]
      ( formats
        [ text_archive ]
        [ binary_archive ]
      )
    ]
  )
| dakota
| ( experimental
[ export_model
  [ filename_prefix STRING ]
  ( formats
    [ text_archive ]
    [ binary_archive ]
  )
]
[ options_file STRING ]
]

| use_derivatives
[ import_build_points_file ALIAS import_points_file STRING
  ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
| annotated
| freeform
| active_only
]
[ export_approx_points_file ALIAS export_points_file STRING
  ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
| annotated
| freeform
]
]
| ea
| lhs ]
[ rng
mt19937
| rnum2
]
[ model_pointer STRING ]
]

| bayes_calibration ALIAS nond_bayes_calibration
  ( queso
    chain_samples ALIAS samples INTEGER
    [ seed INTEGER > 0 ]
    | rng
mt19937
| rnum2
]
5.4 Input Spec Summary

[ emulator
  ( gaussian_process ALIAS kriging
    surfpack
    | dakota
    [ build_samples INTEGER ]
    [ posterior_adaptive ]
    [ import_build_points_file ALIAS import_points_file STRING
      [ { custom.annotated
          [ header ]
          [ eval_id ]
          [ interface_id ]
        }]
      | annotated
      | freeform
      [ active_only ]
    ]
  )
  | pce
  [ p_refinement
    uniform
    |
    ( dimension_adaptive
      sobol
      | decay
      | generalized
    )
  ]
  | [ max_refinement_iterations INTEGER >= 0 ]
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    [ nested
      | non_nested ]
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  | [ sparse_grid_level INTEGER
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      | unrestricted ]
    [ nested
      | non_nested ]
  ]
  | cubature_integrand INTEGER
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    [ dimension_preference REALLIST ]
    [ basis_type
      tensor_product
      | total_order
      |
      adapted
      [ advancements INTEGER ]
      [ soft_convergence_limit INTEGER ]
    ]
  ]
  | collocation_points INTEGER
  [ ( least_squares
    [ svd
      | equality_constrained ]
  )
  ]
  | [ orthogonal_matching_pursuit ALIAS omp
    [ noise_tolerance REALLIST ]
  ]
  | basis.pursuit ALIAS bp
  | { basis.pursuit.denoising ALIAS bpdn
    [ noise_tolerance REALLIST ]
  ]
  | least_angle_regression ALIAS lars
  [ noise_tolerance REALLIST ]
Dakota Input Specification

{ least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]
}

{ cross_validation
  [ noise_only ]
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}

{ ratio_order REAL ]
{ response_scaling ]
{ use_derivatives ]
{ tensor_grid ]
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      [ response_scaling ]
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        [ active_only ]
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      [ posterior_adaptive ]
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| full_covariance ]
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  )
  | rip_sampling ]
[ discrepancy_emulation
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  [ advancements INTEGER ]
  [ soft_convergence_limit INTEGER ]
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[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
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| ( basis_pursuit_denoising ALIAS bpdn
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]
| least_angle_regression ALIAS lars
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| least_absolute_shrinkage ALIAS lasso
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[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
Dakota Input Specification

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5.4 Input Spec Summary

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 | total_order
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 | equality_constrained ]
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 | ( basis_pursuit_denoising ALIAS bpdn
 | [ noise_tolerance REALLIST ]
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 | [ noise_tolerance REALLIST ]
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 | [ least_absolute_shrinkage ALIAS lasso
 | [ noise_tolerance REALLIST ]
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 | [ noise_only ]
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 | [ use_derivatives ]
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\text{\} \text{wiener} \\
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\text{\} \text{p_refinement} \\
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\text{\} local_adaptive} \\
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\text{\} \text{dimension_preference REALLIST} \\
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\text{\} \text{unrestricted} \\
\text{\} \text{nested} \\
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\text{\} \text{use_derivatives} \\
\text{\} \text{diagonal_covariance} \\
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\text{\} mf_sc} \\
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\text{\} \text{dimension_adaptive sobol} \\
\text{\} \text{generalized} \\
\text{\} local_adaptive} \\
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\text{\} \text{unrestricted} \\
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\text{\} \text{non_nested} \} \\
\text{\} \text{piecewise} \\
\text{\} \text{askey} \\
\text{\} \text{wiener} \\
\text{\} \text{use_derivatives} \\
\text{\} \text{diagonal covar}
5.4 Input Spec Summary

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- allocation_control
greedy
- discrepancy_emulation
distinct ALIAS paired
  recursive
- quadrature_order_sequence INTEGERLIST
  dimension_preference REALLIST
  nested
  non_nested
- sparse_grid_level_sequence INTEGERLIST
  dimension_preference REALLIST
  nodal
  hierarchical
  restricted
  unrestricted
  nested
  non_nested
- piecewise
- askey
- wener
- use_derivatives
  diagonal_covariance
  full_covariance
- standardized_space
- logit_transform
- export_chain_points_file STRING
  custom_annotated
  header
  eval_id
  interface_id
  annotated
  freeform
- dram
  delayed_rejection
  adaptive_metropolis
  metropolis_hastings
  multilevel
- pre_solve
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  nip
  none
- proposal_covariance
  prior
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- derivatives
  update_period INTEGER
- values REALLIST
diagonal
  matrix
- filename STRING
diagonal
  matrix
options_file STRING
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5.4 Input Spec Summary

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| sobol |
| decay |
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| non_nested |
|
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| dimension_preference REALIST |
| restricted |
| unrestricted |
| nested |
| non_nested |
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| dimension_preference REALIST |
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| tensor_product |
| total_order |
| ( adapted |
| advancements INTEGER |
| soft_convergence_limit INTEGER |
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| collocation_points INTEGER |
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| svd |
| equality_constrained |
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| ( orthogonal_matching_pursuit ALIAS omp |
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| ( basis_pursuit_denoising ALIAS bpdn |
| noise_tolerance REALIST |
|
| least_angle_regression ALIAS lars |
| noise_tolerance REALIST |
|
| least_absolute_shrinkage ALIAS lasso |
| noise_tolerance REALIST |
| l2_penalty REAL |
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| noise_only |
| max_cv_order_candidates INTEGER >= 0 |
| ratio_order INTEGER |
| response_scaling |
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[ use_derivatives ]
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| equality_constrained ]
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[ noise_tolerance REALLIST ]
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[ noise_tolerance REALLIST ]
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[ noise_tolerance REALLIST ]
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collocation_points INTEGER
[ tensor_grid INTEGERLIST ]
| reuse_points ALIAS reuse_samples ]
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| estimator_variance |
| estimator_rate REAL |
| rip_sampling |
| discrepancy_emulation |
| distinct ALIAS paired |
| recursive |

| expansion_order_sequence INTEGERLIST |
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| basis_type |
| tensor_product |
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| soft_convergence_limit INTEGER |

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| svd |
| equality_constrained |

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| noise_tolerance REALLIST |

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| noise_tolerance REALLIST |

| least_angle_regression ALIAS lars |
| noise_tolerance REALLIST |

| least_absolute_shrinkage ALIAS lasso |
| noise_tolerance REALLIST |
| l2_penalty REAL |

| cross_validation |
| noise_only |
| max_cv_order_candidates INTEGER |

| ratio_order REAL |
| response_scaling |
| use_derivatives |
| tensor_grid |
| reuse_points ALIAS reuse_samples |
| max_solver_iterations INTEGER |

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| reuse_points ALIAS reuse_samples |

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| tensor_grid INTEGERLIST |
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| wiener |
| normalized |
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| full_covariance |
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| uniform |
| ( dimension_adaptive
| sobol |
| decay |
| generalized |
| )
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| greedy |
| discrepancy_emulation
| distinct ALIAS paired |
| recursive |
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| non_nested |
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| dimension_preference REALLIST |
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| dimension_preference REALLIST |
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| tensor_product |
| total_order |
| ( adapted
| advancements INTEGER |
| soft_convergence_limit INTEGER |
)
5.4 Input Spec Summary

[ collocation_ratio REAL
  [ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
  [ ( least_squares
    [ svd
    | equality_constrained ]
  )]

| orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
  |
  | basis_pursuit ALIAS bp
  |
  | basis_pursuit_denoising ALIAS bpdn
  [ noise_tolerance REALLIST ]
  |

| least_angle_regression ALIAS lars
  [ noise_tolerance REALLIST ]
  |

| least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]

| cross_validation
  [ noise_only ]
  [ max_cv_order_candidates INTEGER >= 0 ]

| ratio_order REAL ]
| response_scaling ]
| use_derivatives ]
| tensor_grid ]
| reuse_points ALIAS reuse_samples ]
| max_solver_iterations INTEGER >= 0 ]

| ( expansion_samples_sequence INTEGERLIST
  [ reuse_points ALIAS reuse_samples ]
  ]

| import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )]
  | annotated
  | freeform ]
  | active_only ]

|)

| ( orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
  [ import_build_points_file ALIAS import_points_file STRING
    [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )]
    | annotated
    | freeform ]
    | active_only ]
  ]
  |

| askey
| wiener ]
| normalized ]
| export_expansion_file STRING ]
| diagonal_covariance ]
| full_covariance ]
Dakota Input Specification

```plaintext
|
| sc
|  ( p_refinement
|   uniform
|   ( dimension_adaptive
|    sobol
|    generalized
|   )
| )
|  ( h_refinement
|   uniform
|   ( dimension_adaptive
|    sobol
|    generalized
|   )
|   local_adaptive
| )
| [ max_refinement_iterations INTEGER >= 0 ]
| ( quadrature_order INTEGER
|   ( dimension_preference REALLIST )
|   [ nested
|    non_nested ]
| )
| ( sparse_grid_level INTEGER
|   ( dimension_preference REALLIST )
|   [ nodal
|    hierarchical ]
|   [ restricted
|    unrestricted ]
|   [ nested
|    non_nested ]
| )
| [ piecewise
|  askey
|  wiener ]
| [ use_derivatives ]
| [ diagonal_covariance
|  full_covariance ]
|)
| ( mf_sc
|  ( p_refinement
|   uniform
|   ( dimension_adaptive
|    sobol
|    generalized
|   )
| )
| ( h_refinement
|   uniform
|   ( dimension_adaptive
|    sobol
|    generalized
|   )
|   local_adaptive
| )
| [ max_refinement_iterations INTEGER >= 0 ]
| [ allocation_control
|   greedy ]
| [ discrepancy_emulation
|  distinct ALIAS paired
|   recursive ]
| ( quadrature_order_sequence INTEGERLIST
```

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[ dimension_preference REALLIST ]
| nested
| non_nested }
)

{ sparse_grid_level_sequence INTEGERLIST
| dimension_preference REALLIST ]
| nodal
| hierarchical ]
| restricted
| unrestricted ]
| nested
| non_nested ]
)

[ piecewise
| askey
| wiener ]
| use_derivatives ]
| diagonal_covariance
| full_covariance ]
)

[ standardized_space ]
| data_distribution
| ( gaussian
| means REALLIST
| ( covariance REALLIST
| diagonal
| matrix
|
| obs_data_filename STRING
)
[ posterior_samples_import_filename STRING ]
| generate_posterior_samples
| ( posterior_samples_export_filename STRING ]
)
[ evaluate_posterior_density
| ( posterior_density_export_filename STRING ]
)
)

| dream
| chain_samples ALIAS samples INTEGER
| ( seed INTEGER > 0 ]
| chains INTEGER >= 3 ]
| num_cr INTEGER >= 1 ]
| crossover_chain_pairs INTEGER >= 0 ]
| gr_threshold REAL > 0.0 ]
| jump_step INTEGER >= 0 ]
| emulator
| ( gaussian_process ALIAS kriging
| surfpack
| dakota
| ( build_samples INTEGER ]
| posterior_adaptive ]
| import_build_points_file ALIAS import_points_file STRING
| { custom_annotated
| ( header ]
| eval_id ]
| interface_id ]
)

| annotated
| ( freeform ]
| ( active_only ]
)
)

| ( pce
| ( p_refinement
| uniform
| ( dimension_adaptive
sobol
| decay
| generalized
|
[
  max_refinement_iterations INTEGER >= 0 ]
[
  quadrature_order INTEGER
  [ dimension_preference REALLIST ]
  [ nested
    | non_nested ]
]
[
  sparse_grid_level INTEGER
  [ dimension_preference REALLIST ]
  [ restricted
    | unrestricted ]
  [ nested
    | non_nested ]
]
[
  cubature_integrand INTEGER
]
[
  expansion_order INTEGER
  [ dimension_preference REALLIST ]
  [ basis_type
    tensor_product
    | total_order
    |
    adapted
    [ advancements INTEGER ]
    [ soft_convergence_limit INTEGER ]
  ]
]
[
  collocation_points INTEGER
  [ [ least_squares
    [ svd
    | equality_constrained ]
  ]
]
[
  orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
  [ basis_pursuit ALIAS bp
    |
    [ basis_pursuit_denoising ALIAS bpdn
      [ noise_tolerance REALLIST ]
    ]
    |
    [ least_angle_regression ALIAS lars
      [ noise_tolerance REALLIST ]
    ]
]
[
  least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]
]
[
  cross_validation
  [ noise_only ]
  [ max_cv_order_candidates INTEGER >= 0 ]
]
[
  ratio_order REAL
  [ response_scaling ]
  [ use_derivatives ]
  [ tensor_grid ]
  [ reuse_points ALIAS reuse_samples ]
  [ max_solver_iterations INTEGER >= 0 ]
]
[
  collocation_ratio REAL
  [ [ least_squares
    [ svd
    | equality_constrained ]
  ]
]
|
5.4 Input Spec Summary

{ orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
  }

| basis_pursuit ALIAS bp |
| { basis_pursuit_denoising ALIAS bpdn
  [ noise_tolerance REALLIST ]
  }

| { least_angle_regression ALIAS lars
  [ noise_tolerance REALLIST ]
  }

| { least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ 12_penalty REAL ]
  }

| cross_validation
  [ noise_only ]
  [ max_cv_order_candidates INTEGER >= 0 ]
  }

| ratio_order REAL
| response_scaling
| use_derivatives
| tensor_grid
| { reuse_points ALIAS reuse_samples
  [ max_solver_iterations INTEGER >= 0 ]
  }

| expansion_samples INTEGER
  [ reuse_points ALIAS reuse_samples ]
  }

| import_build_points_file ALIAS import_points_file STRING
  [ { custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
    }
    | annotated
    | freeform
    | active_only
    ]
  [ posterior_adaptive ]
  }

| orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points INTEGER
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
| import_build_points_file ALIAS import_points_file STRING
  [ { custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
    }
    | annotated
    | freeform
    | active_only
    ]
  [ posterior_adaptive ]
  }

| askey
| wiener
| normalized
| export_expansion_file STRING
| diagonal_covariance
| full_covariance
  }

| ml_pce
| allocation_control
  [ estimator_variance
    [ estimator_rate REAL ]
    ]


| rip_sampling |
| discrepancy_emulation
  distinct ALIAS paired
  recursive |
  expansion_order_sequence INTEGERLIST |
  dimension_preference REALLIST |
  basis_type
tensor_product
total_order |
  adapted
  advancements INTEGER |
  soft_convergence_limit INTEGER |
  collocation_ratio REAL |
  collocation_points_sequence ALIAS pilot_samples INTEGERLIST |
  ( least_squares
    svd
    equality_constrained ) |
  orthogonal_matching_pursuit ALIAS omp
    noise_tolerance REALLIST |
  basis_pursuit ALIAS bp |
  basis_pursuit_denoising ALIAS bpdn
    noise_tolerance REALLIST |
  least_angle_regression ALIAS lars
    noise_tolerance REALLIST |
  least_absolute_shrinkage ALIAS lasso
    noise_tolerance REALLIST
    l2_penalty REAL |
  cross_validation
    noise_only
    max_cv_order_candidates INTEGER >= 0 |
  ratio_order REAL |
  response_scaling
    use_derivatives |
  tensor_grid
  reuse_points ALIAS reuse_samples |
  max_solver_iterations INTEGER >= 0 |
  expansion_samples_sequence INTEGERLIST |
  reuse_points ALIAS reuse_samples |
  import_build_points_file ALIAS import_points_file STRING |
  ( custom_annotated
    header
    eval_id
    interface_id ) |
  annotated |
  freeform |
  active_only |
  orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST |
  tensor_grid INTEGERLIST |
  reuse_points ALIAS reuse_samples |
5.4 Input Spec Summary

[ import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  ]
  annotated
  [ freeform ]
  [ active_only ]
]

[ askey
  wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance
  full_covariance ]

[ ( mf_pce
  p_refinement
  uniform ]
  ( dimension_adaptive
    sobol
    decay
    [ generalized
  )]
[ max_refinement_iterations INTEGER >= 0 ]
[ allocation_control
  greedy ]
[ discrepancy_emulation
  distinct ALIAS paired
  recursive ]

[ quadrature_order_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  nested
  non_nested ]

[ sparse_grid_level_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  restricted
  unrestricted
  nested
  non_nested ]

[ expansion_order_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  basis_type
tensor_product
  [ total_order
    [ adapted
      [ advancements INTEGER ]
      [ soft_convergence_limit INTEGER ]
    ]
  ]
  collocation_ratio REAL
[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
[ ( least_squares
  [ svd
    equality_constrained
  ]
  ]
[ orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
  ]
  basis_pursuit ALIAS bp

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basis_pursuit_denoising ALIAS bpdn
[ noise_tolerance REALIST ]

least_angle_regression ALIAS lars
[ noise_tolerance REALIST ]

least_absolute_shrinkage ALIAS lasso
[ noise_tolerance REALIST ]
[ l2_penalty REAL ]

cross_validation
[ noise_only ]
[ max_cv_order_candidates INTEGER >= 0 ]

[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]

expansion_samples_sequence INTEGERLIST
[ reuse_points ALIAS reuse_samples ]

import_build_points_file ALIAS import_points_file STRING
[ { custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
]]
[ annotated ]
[ freeform ]
[ active_only ]

orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
colllocation_points_sequence INTEGERLIST
[ tensor_grid INTEGERLIST ]
[ reuse_points ALIAS reuse_samples ]
import_build_points_file ALIAS import_points_file STRING
[ { custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
]]
[ annotated ]
[ freeform ]
[ active_only ]

askey
wiener
normalized
export_expansion_file STRING
[ diagonal_covariance ]
[ full_covariance ]

sc
[ p_refinement
uniform ]
[ dimension_adaptive
sobol
[ generalized ]
]
5.4 Input Spec Summary

```plaintext
{ h_refinement
  uniform
  |
  ( dimension_adaptive
    sobol
    |
    generalized
  )
  | local_adaptive
  ]
  [ max_refinement_iterations INTEGER >= 0 ]
{ quadrature_order INTEGER
  [ dimension_preference REALLIST ]
  [ nested
    | non_nested ]
  )
  |
{ sparse_grid_level INTEGER
  [ dimension_preference REALLIST ]
  [ nodal
    | hierarchical ]
  [ restricted
    | unrestricted ]
  [ nested
    | non_nested ]
  )
  [ piecewise
    | askey
    | wiener ]
  [ use_derivatives ]
  [ diagonal_covariance
    | full_covariance ]
  )
  |
{ mf_sc
  [ p_refinement
    uniform
    |
    ( dimension_adaptive
      sobol
      |
      generalized
    )
  ]
  |
{ h_refinement
  uniform
  |
  ( dimension_adaptive
    sobol
    |
    generalized
  )
  | local_adaptive
  ]
  [ max_refinement_iterations INTEGER >= 0 ]
  [ allocation_control
    greedy
  ]
  [ discrepancy_emulation
    distinct ALIAS paired
    | recursive
  ]
  [ quadrature_order_sequence INTEGERLIST
    [ dimension_preference REALLIST ]
    [ nested
      | non_nested ]
  ]
  |
{ sparse_grid_level_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  [ nodal
    | hierarchical ]
  [ restricted
    | unrestricted ]
  [ nested
```
non_nested
piecewise
askey
wiener
use_derivatives
diagonal_covariance
full_covariance

standardized_space
export_chain_points_file STRING
{ custom.annotated
  header }
  eval_id
  interface_id
}
annotated
freeform

muq
chain_samples ALIAS samples INTEGER
{ seed INTEGER > 0 }
rng
mt19937
rnum2

dram
delayed_rejection
adaptive_metropolis
metropolis_hastings

proposal_covariance
{ prior
  multiplier REAL > 0.0 }

{ derivatives
  update_period INTEGER }

{ values REALLIST
diagonal
  matrix }

{ filename STRING
diagonal
  matrix }

experimental_design
initial_samples ALIAS samples INTEGER
num_candidates INTEGER > 0
{ max_hifi_evaluations INTEGER >= 0 }
{ batch_size INTEGER >= 1 }
import_candidate_points_file STRING
{ custom.annotated
  header }
  eval_id
  interface_id
}
annotated
freeform

ksg2

calibrate_error_multipliers
one
per_experiment
per_response
5.4 Input Spec Summary

| both |
| [ hyperprior_alphas REALLIST |
|   hyperprior_betas REALLIST ] |
| ] |
| [ burn_in_samples INTEGER ] |
| [ posterior_stats |
|   [ kl_divergence ] |
|   [ mutual_info |
|     [ ksg2 ] |
|   ] |
|   [ kde ] |
| ] |
| [ chain_diagnostics |
|   [ confidence_intervals ] |
| ] |
| [ model_evidence |
|   [ mc_approx ] |
|   [ evidence_samples INTEGER ] |
|   [ laplace_approx ] |
| ] |
| [ model_discrepancy |
|   [ discrepancy_type |
|     ( gaussian_process ALIAS kriging |
|       [ trend_order INTEGER >= 0 ] |
|     ) |
|     ( polynomial |
|       [ basis_order INTEGER >= 0 ] |
|     ) |
| ] |
| [ num_prediction_configs INTEGER >= 0 ] |
| [ prediction_configs REALLIST ] |
| [ import_prediction_configs STRING |
|   [ { custom_annotated |
|     [ header ] |
|     [ eval_id ] |
|     [ interface_id ] |
|   } |
|   annotated |
|   [ freeform ] |
| ] |
| [ export_discrepancy_file STRING |
|   [ { custom_annotated |
|     [ header ] |
|     [ eval_id ] |
|     [ interface_id ] |
|   } |
|   annotated |
|   [ freeform ] |
| ] |
| [ export_corrected_model_file STRING |
|   [ { custom_annotated |
|     [ header ] |
|     [ eval_id ] |
|     [ interface_id ] |
|   } |
|   annotated |
|   [ freeform ] |
| ] |
| [ export_corrected_variance_file STRING |
|   [ { custom_annotated |
|     [ header ] |
|     [ eval_id ] |
|     [ interface_id ] |
|   } |
|   annotated |
|   [ freeform ] |
| ] |
| [ sub_sampling_period INTEGER ] |
| [ probability_levels REALLIST |
|   [ num_probability_levels INTEGERLIST ] |
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ model_pointer STRING ]
[ scaling ]
)
]

dace
grid
| random
| oas
| lhs
| oa_lhs
| box_behnken
| central_composite
| samples INTEGER ]
| seed INTEGER > 0 ]
| fixed_seed ]
| main_effects ]
| quality_metrics ]
| variance_based_decomp
| drop_tolerance REAL ]
]
| symbols INTEGER ]
| model_pointer STRING ]
)

fsu_cvt
| samples INTEGER ]
| seed INTEGER > 0 ]
| fixed_seed ]
| latinize ]
| quality_metrics ]
| variance_based_decomp
| drop_tolerance REAL ]
]
| trial_type
grid
| halton
| random
|
| num_trials INTEGER ]
| max_iterations INTEGER >= 0 ]
| model_pointer STRING ]
)

psuade_moat
| partitions INTEGERLIST ]
| samples INTEGER ]
| seed INTEGER > 0 ]
| model_pointer STRING ]
)

local_evidence ALIAS nond_local_evidence
| sgp
| nip ]
| response_levels REALLIST
| num_response_levels INTEGERLIST ]
| compute
| probabilities
| gen_reliabilities
| system
| series
| parallel
|
|
| probability_levels REALLIST
| num_probability_levels INTEGERLIST ]
|
| gen_reliability_levels REALLIST
| num_gen_reliability_levels INTEGERLIST ]
)
5.4 Input Spec Summary

[ distribution
  cumulative
  | complementary
  ]
[ model_pointer STRING ]
)
)
[ local_interval_est ALIAS nond_local_interval_est
  [ sqp
  | nip ]
  [ convergence_tolerance REAL ]
  [ model_pointer STRING ]
)
)
[ local_reliability ALIAS nond_local_reliability
  [ mpp_search
    x_taylor_mean
    | u_taylor_mean
    | x_taylor_mpp
    | u_taylor_mpp
    | x_two_point
    | u_two_point
    | x_multi_point
    | u_multi_point
    | no_approx
    [ sqp
    | nip ]
  [ integration
    first_order
    | second_order
  [ probability_refinement ALIAS sample_refinement
  import
  | adapt_import
  | mm_adapt_import
  [ refinement_samples INTEGERLIST ]
  [ seed INTEGER > 0 ]
])
]
[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | reliabilities
    | gen_reliabilities
    [ system
      series
      | parallel
      ]
  ]
]
[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  cumulative
  | complementary
  ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ final_moments
  none
  | standard
  | central
  ]
[ model_pointer STRING ]
)
{ global_reliability ALIAS nond_global_reliability
  [ initial_samples INTEGER ]
  x_gaussian_process ALIAS x_kriging
  | u_gaussian_process ALIAS u_kriging
  [ surfpack
   [ export_model
    [ filename_prefix STRING ]
    [ formats
     [ text_archive ]
     [ binary_archive ]
    ]
   ]
  ]
  }

{ experimental
  [ export_model
   [ filename_prefix STRING ]
   [ formats
    [ text_archive ]
    [ binary_archive ]
   ]
  ]
  [ options_file STRING ]
  ]

{ import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  ]
  | annotated
  | freeform
  | active_only
  ]

[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  ]
  | annotated
  | freeform
  ]

[ use_derivatives ]
[ seed INTEGER > 0 ]
[ rng
  mt19937
  | rnum2
  ]

[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | gen_reliabilities
    | system
    series
    | parallel
  ]
  ]

[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]
  ]

[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
  ]

[ distribution
  cumulative
  | complementary
  ]

[ max_iterations INTEGER >= 0 ]
5.4 Input Spec Summary

[ convergence_tolerance REAL ]
[ model_pointer STRING ]
)
|
( fsu_quasi_mc
  | halton
  | hammersley
  [ latinize ]
  [ quality_metrics ]
  [ variance_based_decomp
    [ drop_tolerance REAL ]
  ]
  [ samples INTEGER ]
  [ fixed_sequence ]
  [ sequence_start INTEGERLIST ]
  [ sequence_leap INTEGERLIST ]
  [ prime_base INTEGERLIST ]
  [ max_iterations INTEGER >= 0 ]
  [ model_pointer STRING ]
)
|
( vector_parameter_study
  final_point REALLIST
  | step_vector REALLIST
  num_steps INTEGER
  [ model_pointer STRING ]
)
|
( list_parameter_study
  list_of_points REALLIST
  |
  [ import_points_file STRING
    [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
    | annotated
    | freeform
    | active_only
  ]
  [ model_pointer STRING ]
)
|
( centered_parameter_study
  step_vector REALLIST
  steps_per_variable ALIAS deltas_per_variable INTEGERLIST
  [ model_pointer STRING ]
)
|
( multidim_parameter_study
  partitions INTEGERLIST
  [ model_pointer STRING ]
)
|
( richardson_extrap
  estimate_order
  | converge_order
  | converge_qoi
  [ refinement_rate REAL ]
  [ convergence_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ model_pointer STRING ]
)

KEYWORD model

[ id_model STRING ]
( single ALIAS simulation
  [ interface_pointer STRING ]
  [ solution_level_cost REALLIST
    [ solution_level_control STRING ]
  ]
)
| surrogate
|  [ id_surrogates INTEGERLIST ]
| global
|  ( experimental_gaussian_process
|   | trend
|   | none
|   | constant
|   | linear
|   | reduced_quadratic
|   | quadratic
|   | [ num_restarts INTEGER > 1 ]
|   | [ nugget REAL > 0 ]
|   | [ find_nugget INTEGER ]
|   | [ options_file STRING ]
|   | [ export_approx_variance_file STRING ]
|   | ( custom.annotated
|   |   [ header ]
|   |   [ eval_id ]
|   |   [ interface_id ]
|   |
|   | annotated
|   | freeform
|   |
|   | export_model
|   | [ filename_prefix STRING ]
|   | ( formats
|   |   [ text_archive ]
|   |   [ binary_archive ]
|   |
|   | import_model
|   | [ filename_prefix STRING ]
|   | text_archive
|   | binary_archive
|   |
|   |
|   | ( gaussian_process ALIAS kriging
|   |   ( dakota
|   |   [ point_selection ]
|   |   | trend
|   |   | constant
|   |   | linear
|   |   | reduced_quadratic
|   |
|   |   )
|   |
|   | ( surfpack
|   | trend
|   | constant
|   | linear
|   | reduced_quadratic
|   | quadratic
|   |
|   | optimization_method STRING ]
|   | [ max_trials INTEGER > 0 ]
|   | [ nugget REAL > 0 ]
|   | [ find_nugget INTEGER ]
|   | [ correlation_lengths REALLIST ]
|   | export_model
|   | [ filename_prefix STRING ]
|   | ( formats
|   |   [ text_archive ]
|   |   [ binary_archive ]
|   |   [ algebraic_file ]
|   |
|   | import_model
|   | [ filename_prefix STRING ]
|   | text_archive
5.4 Input Spec Summary

<table>
<thead>
<tr>
<th>binary_archive</th>
</tr>
</thead>
</table>

[ export_approx_variance_file STRING ]

[ ( custom.annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
  )

| annotated |
| freeform |

| mars |
| max_bases INTEGER |

| interpolation linear |

| cubic |

[ export_model |

| filename_prefix STRING ]

( formats
  [ text_archive ]
  [ binary_archive ]
)

[ import_model |

| filename_prefix STRING ]

text_archive |

| binary_archive |

| moving_least_squares |

[ basis_order ALIAS poly_order INTEGER >= 0 ]

[ weight_function INTEGER ]

[ export_model |

| filename_prefix STRING ]

( formats
  [ text_archive ]
  [ binary_archive ]
)

[ import_model |

| filename_prefix STRING ]

text_archive |

| binary_archive |

| function_train |

| regression_type ls |

| rls2 |

| l2_penalty REAL |

| max_solver_iterations INTEGER >= 0 |

| max_cross_iterations INTEGER >= 0 |

| solver_tolerance REAL |

| response_scaling |

| tensor_grid |

| rounding_tolerance REAL |

| arithmetic_tolerance REAL |

| start_order ALIAS order INTEGER >= 0 |

| dimension_preference REALLIST |

| adapt_order |

| kick_order INTEGER > 0 |

| max_order INTEGER >= 0 |

| max_cv_order_candidates INTEGER >= 0 |
[ start_rank ALIAS rank INTEGER >= 0 ]
[ adapt_rank ]
[ kick_rank INTEGER > 0 ]
[ max_rank INTEGER >= 0 ]
[ max_cv_rank_candidates INTEGER >= 0 ]
)

| neural_network
| max_nodes ALIAS nodes INTEGER |
| range REAL |
| random_weight INTEGER |
| export_model
| filename_prefix STRING |
| formats
| text_archive |
| binary_archive |
| algebraic_file |
| algebraic_console |
|
| import_model
| filename_prefix STRING |
| text_archive
| binary_archive |
|
| radial_basis
| bases INTEGER |
| max_pts INTEGER |
| min_partition INTEGER |
| max_subsets INTEGER |
| export_model
| filename_prefix STRING |
| formats
| text_archive |
| binary_archive |
| algebraic_file |
| algebraic_console |
|
| import_model
| filename_prefix STRING |
| text_archive
| binary_archive |
|
| polynomial
| basis_order INTEGER >= 0 |
| linear |
| quadratic |
| cubic |
| export_model
| filename_prefix STRING |
| formats
| text_archive |
| binary_archive |
| algebraic_file |
| algebraic_console |
|
| import_model
| filename_prefix STRING |
| text_archive
| binary_archive |
|
| experimental_polynomial
| basis_order INTEGER >= 0 |
| options_file STRING |
| export_model
5.4 Input Spec Summary

```plaintext
[ filename_prefix STRING ]
[
  formats
  [ text_archive ]
  [ binary_archive ]
]
[
  import_model
  [ filename_prefix STRING ]
  text_archive
  | binary_archive
]
[
  domain_decomposition
  [ cell_type STRING ]
  [ support_layers INTEGER ]
  [ discontinuity_detection
    jump_threshold REAL
    | gradient_threshold REAL
  ]
]
[
  total_points INTEGER
  | minimum_points
  | recommended_points
  [ ( dace_method_pointer STRING
      auto_refinement
      [ max_iterations INTEGER > 0 ]
      [ max_function_evaluations INTEGER > 0 ]
      [ convergence_tolerance REAL ]
      [ soft_convergence_limit INTEGER >= 0 ]
      [ cross_validation_metric STRING
        [ folds INTEGER > 0 ]
      ]
  ]
]
[
  actual_model_pointer ALIAS truth_model_pointer STRING ]
[
  reuse_points ALIAS reuse_samples
  all
  | region
  | none
]
[
  import_build_points_file ALIAS import_points_file ALIAS samples_file STRING
  [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
  ]
  | annotated
  [ use_variable_labels ]
]
  | freeform ]
  | active_only ]
]
[
  export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
  ]
  | annotated
  | freeform ]
]
[
  use_derivatives ]
[
  correction
  [ zeroth_order
    | first_order
    | second_order
  additive
  | multiplicative
  | combined
]
```
[ metrics ALIAS diagnostics STRINGLIST
  [ cross_validation
    [ folds INTEGER
      | percent REAL ]
  ]
  [ press ]
]
[ import_challenge_points_file ALIAS challenge_points_file STRING
  [ ( custom_annotated
    [ header
      [ use_variable_labels ]
    ]
    [ eval_id ]
    [ interface_id ]
  )
  ]
  ( annotated
    [ use_variable_labels ]
    [ freeform ]
    [ active_only ]
  )
]
[ multipoint
  tana
  qmea
  actual_model_pointer ALIAS truth_model_pointer STRING
]
[ local
  taylor_series
  actual_model_pointer ALIAS truth_model_pointer STRING
]
[ hierarchical
  ordered_model_fidelities ALIAS model_fidelity_sequence STRINGLIST
  [ correction
    zeroth_order
    | first_order
    | second_order
    additive
    | multiplicative
    | combined
  ]
]
[ non_hierarchical ALIAS model_ensemble
  truth_model_pointer ALIAS actual_model_pointer STRING
  unordered_model_fidelities ALIAS approximation_models STRINGLIST
]
]
[ nested
  [ optional_interface_pointer STRING
    [ optional_interface_responses_pointer STRING ]
  ]
  [ sub_method_pointer STRING
    [ iterator_servers INTEGER > 0 ]
    [ iterator_scheduling master
      | peer
    ]
    [ processors_per_iterator INTEGER > 0 ]
    [ primary_variable_mapping STRINGLIST ]
    [ secondary_variable_mapping STRINGLIST ]
    [ primary_response_mapping REPLAIST ]
    [ secondary_response_mapping REPLAIST ]
    [ identity_response_mapping ]
  ]
]
[ active_subspace ALIAS subspace
}
5.4 Input Spec Summary

```
actual_model_pointer STRING
[ initial_samples INTEGER ]
[ sample_type
  lhs
  | random
 ]
[ truncation_method
  [ bing_li ]
  [ constantine ]
  [ energy
    [ truncation_tolerance REAL ]
  ]
  [ crossvalidation
    [ minimum
      | relative
      | decrease
    ]
    [ relative_tolerance REAL ]
    [ decrease_tolerance REAL ]
    [ max_rank INTEGER ]
    [ exhaustive ]
  ]
  [ dimension INTEGER ]
  [ bootstrap_samples INTEGER ]
  [ build_surrogate
    [ refinement_samples INTEGERLIST ]
  ]
  [ normalization
    mean_value
    | mean_gradient
    | local_gradient
  ]
]
[ adapted_basis
  [ actual_model_pointer STRING
    sparse_grid_level INTEGER
    |
    ( expansion_order INTEGER
      collocation_ratio REAL
    )
  ]
  [ rotation_method
    unranked
    | ranked
  ]
  [ truncation_tolerance REAL ]
]
[ random_field
  [ build_source
    rf_data_file STRING
    | dace_method_pointer STRING
    |
    ( analytic_covariance
      squared_exponential
      | exponential
    )
  ]
  [ expansion_form
    karhunen_loeve
    | principal_components
  ]
  [ expansion_bases INTEGER ]
  [ truncation_tolerance REAL ]
  [ propagation_model_pointer STRING ]
]
[ variables_pointer STRING ]
[ responses_pointer STRING ]
[ hierarchical_tagging ]
```

KEYWORD: variables
[ id_variables STRING ]
[ active
  all
  | design
  | uncertain
  | aleatory
  | epistemic
  | state
]
[ mixed
  | relaxed ]
[ continuous_design INTEGER > 0
  [ initial_point ALIAS cdv_initial_point REALLIST ]
  [ lower_bounds ALIAS cdv_lower_bounds REALLIST ]
  [ upper_bounds ALIAS cdv_upper_bounds REALLIST ]
  [ scale_types ALIAS cdv_scale_types STRINGLIST ]
  [ scales ALIAS cdv_scales REALLIST ]
  [ descriptors ALIAS cdv_descriptors STRINGLIST ]
]
[ discrete_design_range INTEGER > 0
  [ initial_point ALIAS ddv_initial_point INTEGERLIST ]
  [ lower_bounds ALIAS ddv_lower_bounds INTEGERLIST ]
  [ upper_bounds ALIAS ddv_upper_bounds INTEGERLIST ]
  [ descriptors ALIAS ddv_descriptors STRINGLIST ]
]
[ discrete_design_set
  [ integer INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values INTEGERLIST
    [ categorical STRINGLIST
      [ adjacency_matrix INTEGERLIST ]
    ]
    [ initial_point INTEGERLIST ]
    [ descriptors STRINGLIST ]
  ]
  [ string INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values STRINGLIST
    [ categorical STRINGLIST
      [ adjacency_matrix INTEGERLIST ]
    ]
    [ initial_point STRINGLIST ]
    [ descriptors STRINGLIST ]
  ]
  [ real INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values REALLIST
    [ categorical STRINGLIST
      [ adjacency_matrix INTEGERLIST ]
    ]
    [ initial_point REALLIST ]
    [ descriptors STRINGLIST ]
  ]
]
[ normal_uncertain INTEGER > 0
  means ALIAS nuv_means REALLIST
  std_deviations ALIAS nuv_std_deviations REALLIST
  [ lower_bounds ALIAS nuv_lower_bounds REALLIST ]
  [ upper_bounds ALIAS nuv_upper_bounds REALLIST ]
  [ initial_point REALLIST ]
  [ descriptors ALIAS nuv_descriptors STRINGLIST ]
]
[ lognormal_uncertain INTEGER > 0
  [ lambdas ALIAS lnuv_lambdas REALLIST
    zetas ALIAS lnuv_zetas REALLIST
  ]
  [ means ALIAS lnuv_means REALLIST
    std_deviations ALIAS lnuv_std_deviations REALLIST
    [ error_factors ALIAS lnuv_error_factors REALLIST ]
  ]
  [ lower_bounds ALIAS lnuv_lower_bounds REALLIST ]
  [ upper_bounds ALIAS lnuv_upper_bounds REALLIST ]
  [ initial_point REALLIST ]
  [ descriptors ALIAS lnuv_descriptors STRINGLIST ]
]
uniform_uncertain INTEGER > 0
  lower_bounds ALIAS uuv_lower_bounds REALLIST
  upper_bounds ALIAS uuv_upper_bounds REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS uuv_descriptors STRINGLIST ]
loguniform_uncertain INTEGER > 0
  lower_bounds ALIAS luuv_lower_bounds REALLIST
  upper_bounds ALIAS luuv_upper_bounds REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS luuv_descriptors STRINGLIST ]
triangular_uncertain INTEGER > 0
  modes ALIAS tuv_modes REALLIST
  lower_bounds ALIAS tuv_lower_bounds REALLIST
  upper_bounds ALIAS tuv_upper_bounds REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS tuv_descriptors STRINGLIST ]
exponential_uncertain INTEGER > 0
  betas ALIAS euv_betas REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS euv_descriptors STRINGLIST ]
beta_uncertain INTEGER > 0
  alphas ALIAS buv_alphas REALLIST
  betas ALIAS buv_betas REALLIST
  lower_bounds ALIAS buv_lower_bounds REALLIST
  upper_bounds ALIAS buv_upper_bounds REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS buv_descriptors STRINGLIST ]
gamma_uncertain INTEGER > 0
  alphas ALIAS gauv_alphas REALLIST
  betas ALIAS gauv_betas REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS gauv_descriptors STRINGLIST ]
gumbel_uncertain INTEGER > 0
  alphas ALIAS guuv_alphas REALLIST
  betas ALIAS guuv_betas REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS guuv_descriptors STRINGLIST ]
frechet_uncertain INTEGER > 0
  alphas ALIAS fuv_alphas REALLIST
  betas ALIAS fuv_betas REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS fuv_descriptors STRINGLIST ]
weibull_uncertain INTEGER > 0
  alphas ALIAS wuv_alphas REALLIST
  betas ALIAS wuv_betas REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS wuv_descriptors STRINGLIST ]
histogram_bin_uncertain INTEGER > 0
  [ pairs_per_variable ALIAS num_pairs INTEGERLIST ]
  abscissas ALIAS huv_bin_abscissas REALLIST
  ordinates ALIAS huv_bin_ordinates REALLIST
  counts ALIAS huv_bin_counts REALLIST
  [ initial_point REALLIST ]
  [ descriptors ALIAS huv_bin_descriptors STRINGLIST ]
poisson_uncertain INTEGER > 0
  lambdas REALLIST
  [ initial_point INTEGERLIST ]
  [ descriptors STRINGLIST ]
binomial_uncertain INTEGER > 0
  probability_per_trial ALIAS prob_per_trial REALLIST
  num_trials INTEGERLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]
]

[ negative_binomial_uncertain INTEGER > 0
  probability_per_trial ALIAS prob_per_trial REALLIST
  num_trials INTEGERLIST
  [ initial_point INTEGERLIST ]
  [ descriptors STRINGLIST ]
]

[ geometric_uncertain INTEGER > 0
  probability_per_trial ALIAS prob_per_trial REALLIST
  [ initial_point INTEGERLIST ]
  [ descriptors STRINGLIST ]
]

[ hypergeometric_uncertain INTEGER > 0
  total_population INTEGERLIST
  selected_population INTEGERLIST
  num_drawn INTEGERLIST
  [ initial_point INTEGERLIST ]
  [ descriptors STRINGLIST ]
]

[ histogram_point_uncertain
  [ integer INTEGER > 0
    [ pairs_per_variable ALIAS num_pairs INTEGERLIST
      abscissas INTEGERLIST
      counts REALLIST
      [ initial_point INTEGERLIST ]
      [ descriptors STRINGLIST ]
    ]
  ]

[ string INTEGER > 0
  [ pairs_per_variable ALIAS num_pairs INTEGERLIST
    abscissas STRINGLIST
    counts REALLIST
    [ initial_point STRINGLIST ]
    [ descriptors STRINGLIST ]
  ]

[ real INTEGER > 0
  [ pairs_per_variable ALIAS num_pairs INTEGERLIST
    abscissas REALLIST
    counts REALLIST
    [ initial_point REALLIST ]
    [ descriptors STRINGLIST ]
  ]
]

[ uncertain_correlation_matrix REALLIST ]

[ continuous_interval_uncertain ALIAS interval_uncertain INTEGER > 0
  [ num_intervals ALIAS iuv_num_intervals INTEGERLIST
    [ interval_probabilities ALIAS interval_probs ALIAS iuv_interval_probs REALLIST
      lower_bounds REALLIST
      [ initial_point REALLIST ]
      [ descriptors ALIAS iuv_descriptors STRINGLIST ]
    ]
  ]

[ discrete_interval_uncertain INTEGER > 0
  [ num_intervals INTEGERLIST
    [ interval_probabilities ALIAS interval_probs ALIAS range_probabilities ALIAS range_probs REALLIST
      lower_bounds INTEGERLIST
      [ initial_point INTEGERLIST ]
      [ descriptors STRINGLIST ]
    ]
  ]

[ discrete_uncertain_set
  [ integer INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST
      elements ALIAS set_values INTEGERLIST
      set_probabilities ALIAS set_probs REALLIST
      [ categorical STRINGLIST ]
      [ initial_point INTEGERLIST ]
      [ descriptors STRINGLIST ]
    ]
  ]

[ elements_per_variable ALIAS num_set_values INTEGERLIST
  elements ALIAS set_values STRINGLIST ]
5.4 Input Spec Summary

[ set_probabilities ALIAS set_probs REALLIST ]
[ initial_point STRINGLIST ]
[ descriptors STRINGLIST ]

[ real INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values REALLIST
[ set_probabilities ALIAS set_probs REALLIST ]
categorical STRINGLIST
[ initial_point REALLIST ]
[ descriptors STRINGLIST ]
]

[ continuous_state INTEGER > 0
[ initial_state ALIAS csv_initial_state REALLIST ]
lower_bounds ALIAS csv_lower_bounds REALLIST
upper_bounds ALIAS csv_upper_bounds REALLIST
[ descriptors ALIAS csv_descriptors STRINGLIST ]
]

[ discrete_state_range INTEGER > 0
[ initial_state ALIAS dsv_initial_state INTEGERLIST ]
lower_bounds ALIAS dsv_lower_bounds INTEGERLIST
upper_bounds ALIAS dsv_upper_bounds INTEGERLIST
[ descriptors ALIAS dsv_descriptors STRINGLIST ]
]

[ discrete_state_set
[ integer INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values INTEGERLIST
categorical STRINGLIST
[ initial_state INTEGERLIST ]
[ descriptors STRINGLIST ]
]

[ string INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values STRINGLIST
categorical STRINGLIST
[ initial_state STRINGLIST ]
[ descriptors STRINGLIST ]
]

[ real INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values REALLIST
categorical STRINGLIST
[ initial_state REALLIST ]
[ descriptors STRINGLIST ]
]

[ linear_inequality_constraint_matrix REALLIST ]
[ linear_inequality_lower_bounds REALLIST ]
[ linear_inequality_upper_bounds REALLIST ]
[ linear_inequality_scale_types STRINGLIST ]
[ linear_inequality_scales REALLIST ]
[ linear_equality_constraint_matrix REALLIST ]
[ linear_equality_targets REALLIST ]
[ linear_equality_scale_types STRINGLIST ]
[ linear_equality_scales REALLIST ]

KEYWORD interface
[ id_interface STRING ]
[ analysis_drivers STRINGLIST ]
[ input_filter STRING ]
[ output_filter STRING ]
[ system
[ parameters_file STRING ]
[ results_file STRING ]
[ file_tag ]
[ file_save ]
[ labeled ]
[ aprepro ALIAS dprepro ]
[ work_directory
[ named STRING ]
[ directory_tag ALIAS dir_tag ]
[ directory_save ALIAS dir_save ]
]
[ link_files STRINGLIST ]
[ copy_files STRINGLIST ]
[ replace ]
)
[ allow_existing_results ]
[ verbatim ]
)

(fork
[ parameters_file STRING ]
[ results_file STRING ]
[ file_tag ]
[ file_save ]
[ labeled ]
[ aprepro ALIAS dprepro ]
[ work_directory
[ named STRING ]
[ directory_tag ALIAS dir_tag ]
[ directory_save ALIAS dir_save ]
[ link_files STRINGLIST ]
[ copy_files STRINGLIST ]
[ replace ]
]
[ allow_existing_results ]
[ verbatim ]
)

(direct
[ processors_per_analysis INTEGER > 0 ]
)
| matlab
| pybind11
| python
[ numpy ]
|
| scilab
| grid
[ analysis_components STRINGLIST ]
]
[ algebraic_mappings STRING ]
[ failure_capture
abort
| retry INTEGER
| recover REALLIST
| continuation
]
[ deactivate
[ active_set_vector ]
[ evaluation_cache ]
[ strict_cache_equality
[ cache_tolerance REAL ]
]
[ restart_file ]
]
[ ( batch
[ size INTEGER > 0 ]
)
]

(asynchronous
[ evaluation_concurrency INTEGER > 0 ]
[ local_evaluation_scheduling
dynamic
| static
]
[ analysis_concurrency INTEGER > 0 ]
]
[ evaluation_servers INTEGER > 0 ]
[ evaluation_scheduling
master
| peer
dynamic
]
| static |
| ) |
| processors_per_evaluation INTEGER > 0 |
| analysis_servers INTEGER > 0 |
| analysis_scheduling |
| master |
| peer |

**KEYWORD responses**

```plaintext
| id_responses STRING |
| descriptors ALIAS response_descriptors STRINGLIST |
| objective_functions ALIAS num_objective_functions INTEGER >= 0 |
| sense STRING |
| primary_scale_types ALIAS objective_function_scale_types STRINGLIST |
| primary_scales ALIAS objective_function_scales REALLIST |
| weights ALIAS multi_objective_weights REALLIST |
| nonlinear_inequality_constraints ALIAS num_nonlinear_inequality_constraints INTEGER >= 0 |
| lower_bounds ALIAS nonlinear_inequality_lower_bounds REALLIST |
| upper_bounds ALIAS nonlinear_inequality_upper_bounds REALLIST |
| scale_types ALIAS nonlinear_inequality_scale_types STRINGLIST |
| scales ALIAS nonlinear_inequality_scales REALLIST |
| nonlinear_equality_constraints ALIAS num_nonlinear_equality_constraints INTEGER >= 0 |
| targets ALIAS nonlinear_equality_targets REALLIST |
| scale_types ALIAS nonlinear_equality_scale_types STRINGLIST |
| scales ALIAS nonlinear_equality_scales REALLIST |
| scalar_objectives ALIAS num_scalar_objectives INTEGER >= 0 |
| field_objectives ALIAS num_field_objectives INTEGER >= 0 |
| lengths INTEGERLIST |
| calibration_terms ALIAS least_squares_terms ALIAS num_least_squares_terms INTEGER >= 0 |
| scalar_calibration_terms INTEGER >= 0 |
| field_calibration_terms INTEGER >= 0 |
| lengths INTEGERLIST |
| primary_scales ALIAS calibration_term_scales ALIAS least_squares_term_scales REALLIST |
| weights ALIAS calibration_weights ALIAS least_squares_weights REALLIST |
| num_experiments INTEGER >= 0 |
| num_config_variables INTEGER >= 0 |
| experiment_variance_type ALIAS variance_type STRINGLIST |
| scalar_data_file STRING |
```

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[ nonlinear_inequality_constraints ALIAS num_nonlinear_inequality_constraints INTEGER >= 0 ]
[ lower_bounds ALIAS nonlinear_inequality_lower_bounds REALLIST ]
[ upper_bounds ALIAS nonlinear_inequality_upper_bounds REALLIST ]
[ scale_types ALIAS nonlinear_inequality_scale_types STRINGLIST ]
[ scales ALIAS nonlinear_inequality_scales REALLIST ]
)
[ nonlinear_equality_constraints ALIAS num_nonlinear_equality_constraints INTEGER >= 0 ]
[ targets ALIAS nonlinear_equality_targets REALLIST ]
[ scale_types ALIAS nonlinear_equality_scale_types STRINGLIST ]
[ scales ALIAS nonlinear_equality_scales REALLIST ]
)

( response_functions ALIAS num_response_functions INTEGER >= 0 ]
[ scalar_responses ALIAS num_scalar_responses INTEGER >= 0 ]
[ field_responses ALIAS num_field_responses INTEGER >= 0 ]
[ lengths INTEGERLIST ]
[ num_coordinates_per_field INTEGERLIST ]
[ read_field_coordinates ]
)

no_gradients
| analytic_gradients
| mixed_gradients
| id_numerical_gradients INTEGERLIST
| id_analytic_gradients INTEGERLIST
[ method_source ]
[ ( dakota ]
| ignore_bounds ]
| relative ]
| absolute ]
| bounds ]
| vendor ]
[ interval_type ]
| forward ]
| central ]
[ fd_step_size ALIAS fd_gradient_step_size REALLIST ]
)

( numerical_gradients
[ method_source ]
[ ( dakota ]
| ignore_bounds ]
| relative ]
| absolute ]
| bounds ]
| vendor ]
[ interval_type ]
| forward ]
| central ]
[ fd_step_size ALIAS fd_gradient_step_size REALLIST ]
)

no_hessians

( numerical_hessians
[ fd_step_size ALIAS fd_hessian_step_size REALLIST ]
[ relative ]
| absolute ]
| bounds ]
| forward ]
| central ]
)

| quasi_hessians
| bfgs
| [ damped ]
| srl
| analytic_hessians
mixed_hessians
    [ id_numerical_hessians INTEGERLIST
      [ fd_step_size ALIAS fd_hessian_step_size REALLIST ]
      [ relative
        [ absolute
          [ bounds ]
          [ forward
            [ central ]
        ]
        [ id_quasi_hessians INTEGERLIST
          ( bfgs
            [ damped ]
          )
          [ sr1
            ]
        ]
        [ id_analytic_hessians INTEGERLIST ]
      ]
    )
Chapter 6

Topics Area

This page introduces the user to the topics used to organize keywords.

- admin
- dakota_IO
- dakota_concepts
- models
- variables
- responses
- interface
- methods
- advanced_topics
- packages

6.1 admin

Description

This is only for management while ref man is under construction

Related Topics

- empty
- problem
- not_yet_reviewed

Related Keywords

6.1.1 empty

Description

This topic tracks the keywords which do not have content in the reference manual
Related Topics

Related Keywords

6.1.2 problem

Description

empty

Related Topics

Related Keywords

6.1.3 not_yet_reviewed

Description

Not yet reviewed.

Related Topics

Related Keywords

6.2 dakota_IO

Description

Keywords and Concepts relating inputs to Dakota and outputs from Dakota

Related Topics

• dakota_inputs
• dakota_output
• file_formats

Related Keywords

• error_file : Base filename for error redirection
• output_file : Base filename for output redirection
• input : Base filename for post-run mode data input
• output : Base filename for post-run mode data output
• input : Base filename for pre-run mode data input
• output : Base filename for pre-run mode data output
• read_restart : Base filename for restart file read
• stop_restart : Evaluation ID number at which to stop reading restart file
6.2 dakota_IO

- **input**: Base filename for run mode data input
- **output**: Base filename for run mode data output
- **write_restart**: Base filename for restart file write

6.2.1 dakota_inputs

**Description**

`empty`

**Related Topics**

- **block**
- **data_import_capabilities**

**Related Keywords**

6.2.1.1 block

**Description**

A block is the highest level of keyword organization in Dakota. There are currently 6 blocks in the Dakota input spec:

**Related Topics**

- **block_identifier**
- **block_pointer**

**Related Keywords**

- **environment**: Top-level settings for Dakota execution
- **interface**: Specifies how function evaluations will be performed in order to map the variables into the responses.
- **method**: Begins Dakota method selection and behavioral settings.
- **model**: Specifies how variables are mapped into a set of responses
- **responses**: Description of the model output data returned to Dakota upon evaluation of an interface.
- **variables**: Specifies the parameter set to be iterated by a particular method.

6.2.1.2 block_identifier

**Description**

`empty`
Related Topics

Related Keywords

- **id_interface**: Name the interface block; helpful when there are multiple
- **id_method**: Name the method block; helpful when there are multiple
- **id_model**: Give the model block an identifying name, in case of multiple model blocks
- **id_responses**: Name the responses block; helpful when there are multiple
- **id_variables**: Name the variables block; helpful when there are multiple

6.2.1.3 block_pointer

Description

See block_pointer for details about pointers.

Related Topics

Related Keywords

- **top_method_pointer**: Identify which method leads the Dakota study
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
- **method_pointer**: Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
- **model_pointer**: Identifier for model block to be used by a method
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- `model_pointer`: Identifier for model block to be used by a method
- `model_pointer`: Identifier for model block to be used by a method
- `model_pointer_list`: Associate models with method names
- `method_pointer_list`: Pointers to methods to execute sequentially or collaboratively
- `global_model_pointer`: Pointer to model used by global method
- `global_method_pointer`: Pointer to global method
- `local_model_pointer`: Pointer to model used by local method
- `local_method_pointer`: Pointer to local method
- `model_pointer_list`: Associate models with method names
- `method_pointer_list`: Pointers to methods to execute sequentially or collaboratively
- `model_pointer`: Identifier for model block to be used by a method
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- `model_pointer`: Identifier for model block to be used by a method
- `model_pointer`: Identifier for model block to be used by a method
- `method_pointer`: Pointer to sub-method to run from each starting point
- `model_pointer`: Identifier for model block to be used by a method
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• **model_pointer**: Identifier for model block to be used by a method
• **model_pointer**: Identifier for model block to be used by a method
• **method_pointer**: Pointer to optimization or least-squares sub-method
• **model_pointer**: Identifier for model block to be used by a method
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• **model_pointer**: Identifier for model block to be used by a method
• **model_pointer**: Identifier for model block to be used by a method
• **actual_model_pointer**: Pointer to specify a full-space model, from which to construct a lower dimensional surrogate
• **actual_model_pointer**: Pointer to specify a “truth” model, from which to construct a surrogate
• **optional_interface_pointer**: Pointer to interface that provides non-nested responses

• **optional_interface_responses_pointer**: Pointer to responses block that defines non-nested responses

• **sub_method_pointer**: The sub_method_pointer specifies the method block for the sub-iterator

• **responses_pointer**: Specify which responses block will be used by this model block

• **interface_pointer**: Interface block pointer for the single model type

• **dace_method_pointer**: Specify a method to gather training data

• **actual_model_pointer**: Pointer to specify a “truth” model, from which to construct a surrogate

• **truth_model_pointer**: Pointer to specify a “truth” model, from which to construct a surrogate

• **variables_pointer**: Specify which variables block will be included with this model block

### 6.2.1.4 data_import_capabilities

**Description**

empty

**Related Topics**

**Related Keywords**

### 6.2.2 dakota_output

**Description**

empty

**Related Topics**

**Related Keywords**

• **graphics**: (DEPRECATED) Display plots of variables and responses

• **output_precision**: Control the output precision

• **results_output**: (Experimental) Write a summary file containing the final results

• **hdf5**: Write results to file in HDF5 format

• **interface_selection**: Select the models that write evaluation data to HDF5

• **all**: Write evaluation data for all interfaces to HDF5

• **none**: Write evaluation data for no interfaces to HDF5

• **simulation**: Write evaluation data only for simulation interfaces to HDF5

• **model_selection**: Select the models that write evaluation data to HDF5

• **all**: Write evaluation data to HDF5 for all models

• **all_methods**: Write evaluation data to HDF5 for all models that belong directly to methods
• **none**: Write evaluation data for no models to HDF5
• **top_method**: Write evaluation data only for the top-level method's model to HDF5
• **results_output_file**: The base file name of the results file
• **text**: Write results to file in text format
• **tabular_data**: Write a tabular results file with variable and response history
• **tabular_data_file**: File name for tabular data output
• **output**: Control how much method information is written to the screen and output file

### 6.2.3 file_formats

**Description**

See sections “Inputs to Dakota” and “Outputs from Dakota” in the Dakota User’s Manual[5].

**Related Topics**

**Related Keywords**

• **annotated**: Selects annotated tabular file format
• **custom_annotated**: Selects custom-annotated tabular file format
• **freeform**: Selects freeform file format
• **annotated**: Selects annotated tabular file format
• **custom_annotated**: Selects custom-annotated tabular file format
• **freeform**: Selects freeform file format
• **annotated**: Selects annotated tabular file format
• **custom_annotated**: Selects custom-annotated tabular file format
• **freeform**: Selects freeform file format
• **aprepro**: Write parameters files in APREPRO syntax
• **labeled**: Requires correct function value labels in results file
• **aprepro**: Write parameters files in APREPRO syntax
• **labeled**: Requires correct function value labels in results file
• **annotated**: Selects annotated tabular file format
• **custom_annotated**: Selects custom-annotated tabular file format
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• **annotated** : Selects annotated tabular file format
• **custom_annotated** : Selects custom-annotated tabular file format
• **freeform** : Selects freeform file format
• **active_only** : Import only active variables from tabular data file
• **annotated** : Selects annotated tabular file format
• **custom_annotated** : Selects custom-annotated tabular file format
• **freeform** : Selects freeform file format
• **active_only** : Import only active variables from tabular data file
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• **custom_annotated** : Selects custom-annotated tabular file format
• **freeform** : Selects freeform file format
• **active_only** : Import only active variables from tabular data file
6.3 dakota_concepts

Description

Miscellaneous concepts related to Dakota operation
6.3 dakota_concepts

Related Topics

- method_independent_controls
- block
- strategies
- command_line_options
- restarts
- pointers

Related Keywords

6.3.1 method_independent_controls

Description

The <method independent controls> are those controls which are valid for a variety of methods. In some cases, these controls are abstractions which may have slightly different implementations from one method to the next. While each of these controls is not valid for every method, the controls are valid for enough methods that it was reasonable to consolidate the specifications.

Related Topics

Related Keywords

- max_iterations : Number of iterations allowed for optimizers and adaptive UQ methods
- convergence_tolerance : Stopping criterion based on relative error reduction
- max_function_evaluations : Stopping criterion based on maximum function evaluations
- max_iterations : Number of iterations allowed for optimizers and adaptive UQ methods
- constraint_tolerance : Maximum allowable constraint violation still considered feasible
- max_function_evaluations : Number of function evaluations allowed for optimizers
- scaling : Turn on scaling for variables, responses, and constraints
- convergence_tolerance : Stopping criterion based on objective function or statistics convergence
- max_iterations : Number of iterations allowed for optimizers and adaptive UQ methods
- scaling : Turn on scaling for variables, responses, and constraints
- scaling : Turn on scaling for variables, responses, and constraints
- convergence_tolerance : Stopping criterion based on objective function or statistics convergence
- max_function_evaluations : Number of function evaluations allowed for optimizers
- max_iterations : Number of iterations allowed for optimizers and adaptive UQ methods
- scaling : Turn on scaling for variables, responses, and constraints
- convergence_tolerance : Stopping criterion based on objective function or statistics convergence
- max_function_evaluations : Number of function evaluations allowed for optimizers
- max_iterations : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **speculative**: Compute speculative gradients

• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **speculative**: Compute speculative gradients

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

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• **scaling**: Turn on scaling for variables, responses, and constraints

• **speculative**: Compute speculative gradients

• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **scaling**: Turn on scaling for variables, responses, and constraints

• **speculative**: Compute speculative gradients

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **final_solutions**: Number of designs returned as the best solutions

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods

• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence

• **max_function_evaluations**: Number of function evaluations allowed for optimizers

• **scaling**: Turn on scaling for variables, responses, and constraints

• **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **scaling**: Turn on scaling for variables, responses, and constraints
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **id_method**: Name the method block; helpful when there are multiple
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **scaling**: Turn on scaling for variables, responses, and constraints
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **scaling**: Turn on scaling for variables, responses, and constraints
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
• **max_iterations** : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling** : Turn on scaling for variables, responses, and constraints
• **convergence_tolerance** : Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations** : Number of function evaluations allowed for optimizers
• **max_iterations** : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling** : Turn on scaling for variables, responses, and constraints
• **speculative** : Compute speculative gradients
• **convergence_tolerance** : Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations** : Number of function evaluations allowed for optimizers
• **max_iterations** : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling** : Turn on scaling for variables, responses, and constraints
• **speculative** : Compute speculative gradients
• **convergence_tolerance** : Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations** : Number of function evaluations allowed for optimizers
• **max_iterations** : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling** : Turn on scaling for variables, responses, and constraints
• **speculative** : Compute speculative gradients
• **convergence_tolerance** : Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations** : Number of function evaluations allowed for optimizers
• **max_iterations** : Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling** : Turn on scaling for variables, responses, and constraints
• **speculative** : Compute speculative gradients
• **convergence_tolerance** : Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations**: Number of function evaluations allowed for optimizers
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **speculative**: Compute speculative gradients
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations**: Number of function evaluations allowed for optimizers
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **speculative**: Compute speculative gradients
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
• **max_function_evaluations**: Number of function evaluations allowed for optimizers
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **speculative**: Compute speculative gradients
• **output**: Control how much method information is written to the screen and output file
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
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• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
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• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
• **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
• **constraint_tolerance**: Maximum allowable constraint violation still considered feasible
• **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
• **scaling**: Turn on scaling for variables, responses, and constraints
6.3.2 block

Description

A block is the highest level of keyword organization in Dakota. There are currently 6 blocks in the Dakota input spec:

Related Topics

- block_identifier
- block_pointer

Related Keywords

- environment: Top-level settings for Dakota execution
- interface: Specifies how function evaluations will be performed in order to map the variables into the responses.
- model: Specifies how variables are mapped into a set of responses
- responses: Description of the model output data returned to Dakota upon evaluation of an interface.
- variables: Specifies the parameter set to be iterated by a particular method.

6.3.2.1 block_identifier

Description

empty

Related Topics

Related Keywords

- id_interface: Name the interface block; helpful when there are multiple
- id_method: Name the method block; helpful when there are multiple
- id_model: Give the model block an identifying name, in case of multiple model blocks
- id_responses: Name the responses block; helpful when there are multiple
- id_variables: Name the variables block; helpful when there are multiple

6.3.2.2 block_pointer

Description

See block_pointer for details about pointers.
Related Topics

Related Keywords

- top_method_pointer: Identify which method leads the Dakota study
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- method_pointer: Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
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- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
• **model_pointer_list**: Associate models with method names
• **method_pointer_list**: Pointers to methods to execute sequentially or collaboratively
• **global_model_pointer**: Pointer to model used by global method
• **global_method_pointer**: Pointer to global method
• **local_model_pointer**: Pointer to model used by local method
• **local_method_pointer**: Pointer to local method
• **model_pointer_list**: Associate models with method names
• **method_pointer_list**: Pointers to methods to execute sequentially or collaboratively
• **model_pointer**: Identifier for model block to be used by a method
• **method_pointer**: Pointer to sub-method to run from each starting point
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• method_pointer: Pointer to optimization or least-squares sub-method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• method_pointer: Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
• model_pointer: Identifier for model block to be used by a method
• method_pointer: Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• model_pointer: Identifier for model block to be used by a method
• actual_model_pointer: Pointer to specify a full-space model, from which to construct a lower dimensional surrogate
• actual_model_pointer: Pointer to specify a "truth" model, from which to construct a surrogate
• optional_interface_pointer: Pointer to interface that provides non-nested responses
• optional_interface_responses_pointer: Pointer to responses block that defines non-nested responses
• sub_method_pointer: The sub_method_pointer specifies the method block for the sub-iterator
• responses_pointer: Specify which responses block will be used by this model block
• interface_pointer: Interface block pointer for the single model type
• dace_method_pointer: Specify a method to gather training data
• actual_model_pointer: Pointer to specify a "truth" model, from which to construct a surrogate
• truth_model_pointer: Pointer to specify a "truth" model, from which to construct a surrogate
• variables_pointer: Specify which variables block will be included with this model block
6.3.3 strategies

Description
empty

Related Topics

• advanced_strategies

Related Keywords

6.3.3.1 advanced_strategies

Description
empty

Related Topics

Related Keywords

6.3.4 command_line_options

Description
empty

Related Topics

Related Keywords

• check : Invoke Dakota in input check mode
• error_file : Base filename for error redirection
• output_file : Base filename for output redirection
• post_run : Invoke Dakota with post-run mode active
• pre_run : Invoke Dakota with pre-run mode active
• read_restart : Base filename for restart file read
• run : Invoke Dakota with run mode active
• write_restart : Base filename for restart file write

6.3.5 restarts

Description
empty
6.3.6 pointers

Description

For all pointer specifications, if a pointer string is specified and no corresponding id string is available, Dakota will exit with an error message.

If the pointer is optional and no pointer string is specified, then the last specification parsed will be used.

It is appropriate to omit optional cross-referencing whenever the relationships are unambiguous due to the presence of only one specification.

Related Topics

- block_pointer
- objective_function_pointer

Related Keywords

6.3.6.1 block_pointer

Description

See block_pointer for details about pointers.

Related Topics

Related Keywords

- top_method_pointer: Identify which method leads the Dakota study
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
- model_pointer: Identifier for model block to be used by a method
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• `model_pointer` : Identifier for model block to be used by a method

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• `model_pointer` : Identifier for model block to be used by a method

• `method_pointer_list` : Pointers to methods to execute sequentially or collaboratively

• `global_model_pointer` : Pointer to model used by global method

• `global_method_pointer` : Pointer to global method

• `local_model_pointer` : Pointer to model used by local method

• `local_method_pointer` : Pointer to local method

• `model_pointer_list` : Associate models with method names

• `method_pointer_list` : Pointers to methods to execute sequentially or collaboratively

• `model_pointer` : Identifier for model block to be used by a method

• `model_pointer` : Identifier for model block to be used by a method

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• `model_pointer` : Identifier for model block to be used by a method

• `model_pointer` : Identifier for model block to be used by a method

• `model_pointer` : Identifier for model block to be used by a method

• `method_pointer` : Pointer to sub-method to run from each starting point
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
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• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **method_pointer** : Pointer to optimization or least-squares sub-method
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
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• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **model_pointer** : Identifier for model block to be used by a method
• **method_pointer** : Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
• **model_pointer** : Identifier for model block to be used by a method
6.4 models

- `method_pointer` : Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem
- `model_pointer` : Identifier for model block to be used by a method
- `model_pointer` : Identifier for model block to be used by a method
- `model_pointer` : Identifier for model block to be used by a method
- `actual_model_pointer` : Pointer to specify a full-space model, from which to construct a lower dimensional surrogate
- `actual_model_pointer` : Pointer to specify a “truth” model, from which to construct a surrogate
- `optional_interface_pointer` : Pointer to interface that provides non-nested responses
- `optional_interface_responses_pointer` : Pointer to responses block that defines non-nested responses
- `sub_method_pointer` : The sub_method_pointer specifies the method block for the sub-iterator
- `responses_pointer` : Specify which responses block will be used by this model block
- `interface_pointer` : Interface block pointer for the single model type
- `dace_method_pointer` : Specify a method to gather training data
- `actual_model_pointer` : Pointer to specify a “truth” model, from which to construct a surrogate
- `actual_model_pointer` : Pointer to specify a “truth” model, from which to construct a surrogate
- `truth_model_pointer` : Pointer to specify a “truth” model, from which to construct a surrogate
- `variables_pointer` : Specify which variables block will be included with this model block

6.3.6.2 objective_function_pointer

Description

See `block_pointer` for details about pointers.

Related Topics

Related Keywords

- `id_analytic_gradiens` : Identify which analytical gradient corresponds to which response
- `id_numerical_gradients` : Identify which numerical gradient corresponds to which response
- `id_analytic_hessians` : Identify which analytical Hessian corresponds to which response
- `id_numerical_hessians` : Identify which numerical-Hessian corresponds to which response
- `id_quasi_hessians` : Identify which quasi-Hessian corresponds to which response

6.4 models

Description

Keywords and Concepts relating to the `model` block
Related Topics

- surrogate_models
- recast_models
- multifidelity_models
- reduced_order_models
- nested_models
- advanced_model_recursion

Related Keywords

### 6.4.1 surrogate_models

Description

empty

Related Topics

- surrogate_based_optimization_methods

Related Keywords

- export_model: Exports surrogate model in user-specified format(s)
- filename_prefix: User-customizable portion of exported/imported surrogate model filenames
- formats: Formats for surrogate model export
- binary_archive: Surrogate model binary archive file format
- text_archive: Surrogate model plain-text archive file format
- export_model: Exports surrogate model in user-specified format(s)
- filename_prefix: User-customizable portion of exported/imported surrogate model filenames
- formats: Formats for surrogate model export
- binary_archive: Surrogate model binary archive file format
- text_archive: Surrogate model plain-text archive file format
- export_model: Exports surrogate model in user-specified format(s)
- filename_prefix: User-customizable portion of exported/imported surrogate model filenames
- formats: Formats for surrogate model export
- binary_archive: Surrogate model binary archive file format
- text_archive: Surrogate model plain-text archive file format
- export_model: Exports surrogate model in user-specified format(s)
- filename_prefix: User-customizable portion of exported/imported surrogate model filenames
- formats: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **auto_refinement**: Experimental auto-refinement of surrogate model
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **point_selection**: Enable greedy selection of well-spaced build points
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **algebraic_console**: Export surrogate model in algebraic format to the console
• **algebraic_file**: Export surrogate model in algebraic format to a file
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **metrics**: Compute surrogate quality metrics
• **cross_validation**: Perform k-fold cross validation
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **export_model**: Exports surrogate model in user-specified format(s)
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **formats**: Formats for surrogate model export
• **algebraic_console**: Export surrogate model in algebraic format to the console
• **algebraic_file**: Export surrogate model in algebraic format to a file
• **binary_archive**: Surrogate model binary archive file format
• **text_archive**: Surrogate model plain-text archive file format
• **binary_archive**: Surrogate model binary archive file format
• **filename_prefix**: User-customizable portion of exported/imported surrogate model filenames
• **text_archive**: Surrogate model plain-text archive file format
• **max_nodes**: Maximum number of hidden layer nodes
• **random_weight**: (Inactive) Random weight control
• **range**: Range for neural network random weights
• **export_model**: Exports surrogate model in user-specified format(s)
• *filename_prefix*: User-customizable portion of exported/imported surrogate model filenames

• *formats*: Formats for surrogate model export

• *algebraic_console*: Export surrogate model in algebraic format to the console

• *algebraic_file*: Export surrogate model in algebraic format to a file

• *binary_archive*: Surrogate model binary archive file format

• *text_archive*: Surrogate model plain-text archive file format

• *binary_archive*: Surrogate model binary archive file format

• *filename_prefix*: User-customizable portion of exported/imported surrogate model filenames

• *text_archive*: Surrogate model plain-text archive file format

• *bases*: Initial number of radial basis functions

• *export_model*: Exports surrogate model in user-specified format(s)

• *filename_prefix*: User-customizable portion of exported/imported surrogate model filenames

• *formats*: Formats for surrogate model export

• *algebraic_console*: Export surrogate model in algebraic format to the console

• *algebraic_file*: Export surrogate model in algebraic format to a file

• *binary_archive*: Surrogate model binary archive file format

• *text_archive*: Surrogate model plain-text archive file format

• *binary_archive*: Surrogate model binary archive file format

• *filename_prefix*: User-customizable portion of exported/imported surrogate model filenames

• *text_archive*: Surrogate model plain-text archive file format

• *max_pts*: Maximum number of RBF CVT points

• *max_subsets*: Number of trial RBF subsets

• *min_partition*: (Inactive) Minimum RBF partition

• *reuse_points*: Surrogate model training data reuse control

### 6.4.2 surrogate_based_optimization_methods

**Description**

empty

**Related Topics**

**Related Keywords**

• *efficient_global*: Global Surrogate Based Optimization, a.k.a. EGO

• *surrogate_based_global*: Global Surrogate Based Optimization

• *surrogate_based_local*: Local Surrogate Based Optimization
6.4.3 recast_models

Description
empty

Related Topics
Related Keywords

6.4.4 multifidelity_models

Description
empty

Related Topics
Related Keywords

6.4.5 reduced_order_models

Description
empty

Related Topics
Related Keywords

6.4.6 nested_models

Description
empty

Related Topics
Related Keywords

6.4.7 advanced_model_recursion

Description
empty

Related Topics

• hybrid_and_recursions_logic
6.5 variables

Related Keywords

6.4.8 hybrid_and_recursions_logic

Description

empty

Related Topics

Related Keywords

6.5 variables

Description

Keywords and concepts relating to the variables block, which defines the parameters for the study.

Related Topics

- variable_domain
- linear_constraints
- variable_type

Related Keywords

6.5.1 variable_domain

Description

Dakota variables can be grouped by their valid domains.

1. Mixed: continuous and discrete variables are treated separately
2. Relaxed: noncategorical discrete variables are relaxed and treated as continuous variables (categorical variables are non-relaxable and remain discrete)

Refer to mixed and relaxed for additional information.

Related Topics

- continuous_variables
- discrete_variables

Related Keywords

6.5.2 continuous_variables

Description

This page collects information related to the topic of continuous design, uncertain, and state variables.
Related Keywords

- **beta_uncertain**: Aleatory uncertain variable - beta
- **continuous_design**: Design variable - continuous
- **continuous_interval_uncertain**: Epistemic uncertain variable - values from one or more continuous intervals
- **continuous_state**: State variable - continuous
- **exponential_uncertain**: Aleatory uncertain variable - exponential
- **frechet_uncertain**: Aleatory uncertain variable - Frechet
- **gamma_uncertain**: Aleatory uncertain variable - gamma
- **gumbel_uncertain**: Aleatory uncertain variable - gumbel
- **histogram_bin_uncertain**: Aleatory uncertain variable - continuous histogram
- **lognormal_uncertain**: Aleatory uncertain variable - lognormal
- **loguniform_uncertain**: Aleatory uncertain variable - loguniform
- **normal_uncertain**: Aleatory uncertain variable - normal (Gaussian)
- **triangular_uncertain**: Aleatory uncertain variable - triangular
- **uniform_uncertain**: Aleatory uncertain variable - uniform
- **weibull_uncertain**: Aleatory uncertain variable - Weibull

6.5.3 discrete_variables

Description

This page discusses discrete design, uncertain, and state variables (which have discrete in their keyword name) as they have similar specifications. These include:

1. Integer ranges
2. Sets of integers
3. Sets of reals
4. Sets of strings and each is described below.

In addition, some aleatory uncertain variables, e.g., **binomial_uncertain**, are discrete integer-valued random variables specified using parameters. These are described on their individual keyword pages.

Sets

Sets of integers, reals, and strings have similar specifications, though different value types.

The variables are specified using three keywords:

- **Variable declaration keyword** - specifies the number of variables being defined
- **elements_per_variable** - a list of positive integers specifying how many set members each variable admits
  - Length = # of variables
6.5 variables

- **elements** - a list of the permissible integer values in ALL sets, concatenated together.
  - Length = sum of elements_per_variable, or an integer multiple of number of variables
  - The order is very important here.
  - The list is partitioned according to the values of elements_per_variable, and each partition is assigned to a variable.

- The ordering of elements_per_variable, and the partitions of elements must match the strings from descriptors

For string variables, each string element value must be quoted and may contain alphanumeric, dash, underscore, and colon. White space, quote characters, and backslash/metacharacters are not permitted.

Examples are given on the pages:

- discrete design set **integer**
- discrete design set **real**
- discrete design set **string**
- discrete uncertain set **integer**
- discrete uncertain set **real**
- discrete uncertain set **string**

**Range**

For discrete variables defined by range(s), the **lower_bounds** and **upper_bounds** restrict the permissible values. For design variables, this constrains the feasible design space and is frequently used to prevent nonphysical designs. This is a discrete interval variable that may take any integer value within bounds (e.g., [1, 4], allowing values of 1, 2, 3, or 4). For some variable types, each variable is can be defined by multiple ranges.

Examples are given on the pages:

- discrete_interval_uncertain

**Related Topics**

**Related Keywords**

- **binomial_uncertain** : Aleatory uncertain discrete variable - binomial
- **discrete_design_range** : Design variable - discrete range-valued
- **discrete_design_set** : Design variable - discrete set-valued
- **integer** : Integer-valued discrete design variables
- **real** : Real-valued discrete design variables
- **string** : String-valued discrete design set variables
- **discrete_interval_uncertain** : Epistemic uncertain variable - values from one or more discrete intervals
- **discrete_state_range** : State variables - discrete range-valued
- **discrete_state_set** : State variable - discrete set-valued
- **integer** : Discrete state variables, each defined by a set of permissible integers
- **real** : Discrete state variables, each defined by a set of permissible real numbers
• **string**: String-valued discrete state set variables
• **discrete_uncertain_set**: Epistemic uncertain variable - discrete set-valued
• **integer**: Discrete, epistemic uncertain variable - integers within a set
• **real**: Discrete, epistemic uncertain variable - real numbers within a set
• **string**: Discrete, epistemic uncertain variable - strings within a set
• **geometric_uncertain**: Aleatory uncertain discrete variable - geometric
• **histogram_point_uncertain**: Aleatory uncertain variable - discrete histogram
• **hypergeometric_uncertain**: Aleatory uncertain discrete variable - hypergeometric
• **negative_binomial_uncertain**: Aleatory uncertain discrete variable - negative binomial
• **poisson_uncertain**: Aleatory uncertain discrete variable - Poisson

### 6.5.4 linear_constraints

**Description**

Many methods can make use of linear equality or inequality constraints. As the name implies, linear constraints are constraints that are linear functions of the variables. Constraints that are nonlinear functions of variables are specified using the **nonlinear_constraints** family of keywords. From a Dakota usage point of view, the most important difference between linear and nonlinear constraints is that the former are specified entirely within the Dakota input file and calculated by Dakota itself, while the latter must be calculated by the user's simulation and returned as responses to Dakota.

The Optimization chapter of the User's Manual[5] states which methods support linear constraints. Of those methods, a subset strictly obey linear constraints; that is, no candidate points are generated by the optimizer that violate the constraints. These include asynch_pattern_search, the optpp_* family of optimizers (with the exception of optpp_fd_newton), and npsol_sqp. The other methods seek feasible solutions (i.e. solutions that satisfy the linear constraints), but may violate the constraints as they run. Linear constraints may also be violated, even when using an optimizer that itself strictly respects them, if numerical_gradients are used. In this case, Dakota may request evaluations that lie outside of the feasible region when computing a gradient near the boundary.

One final limitation that bears mentioning is that linear constraints are compatible only with continuous variables. No discrete types are permitted when using linear constraints.

**Related Topics**

**Related Keywords**

- **linear_equation_constraint_matrix**: Define coefficients of the linear equalities
- **linear_equation_scale_types**: How to scale each linear equality constraint
- **linear_equation_scales**: Characteristic values to scale linear equalities
- **linear_equation_targets**: Define target values for the linear equality constraints
- **linear_inequality_constraint_matrix**: Define coefficients of the linear inequality constraints
- **linear_inequality_lower_bounds**: Define lower bounds for the linear inequality constraint
- **linear_inequality_scale_types**: How to scale each linear inequality constraint
- **linear_inequality_scales**: Characteristic values to scale linear inequalities
- **linear_inequality_upper_bounds**: Define upper bounds for the linear inequality constraint

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6.5.5 variable_type

Description

Dakota variables can be grouped their type: design, uncertain (further sub-divided into aleatory and epistemic), and state. These distinctions are useful in situations where the user may want to explicitly control the subset of variables that is considered active for a certain Dakota method, overriding the default active variable type.

Related Topics

- design_variables
- aleatory_uncertain_variables
- epistemic_uncertain_variables
- state_variables

Related Keywords

6.5.6 design_variables

Description

Design variables are adjusted in the course of seeking an optimal design or an optimal set of deterministic calibration parameters.

Continuous design variables, which may assume any real value within specified bounds, are the most common design variable type in engineering applications. All but a handful of the optimization algorithms in Dakota support continuous design variables exclusively.

Related Topics

Related Keywords

6.5.7 aleatory_uncertain_variables

Description

Aleatory uncertainty is also known as inherent variability, irreducible uncertainty, or randomness.

Aleatory uncertainty is typically represented by probability distributions and is specified to Dakota using the following parametric and histogram uncertain variable keywords.
6.5.8 epistemic_uncertain_variables

Description

Epistemic uncertainty is uncertainty due to lack of knowledge.

In Dakota, epistemic uncertainty can be characterized by interval- or set-valued variables (see relevant keywords below) that are propagated to calculate bounding intervals on simulation output using interval analysis methods. These epistemic variable types can optionally include basic probability assignments for use in Dempster-Shafer theory of evidence methods. Epistemic uncertainty can alternately be modeled with probability density functions, although results from UQ studies are then typically interpreted as possibilities or bounds, as opposed to a probability distribution of responses.

Through nested models, Dakota can perform combined aleatory / epistemic analyses such as second-order probability or probability of frequency. For example, a variable can be assumed to have a lognormal distribution with specified variance, with its mean expressed as an epistemic uncertainty lying in an expert-specified interval.
6.6 responses

Description

Keywords and concepts relating to the responses block

Related Topics

- response_types
- nonlinearConstraints
Related Keywords

6.6.1 response_types

Description

The specification must be one of three types:

1. objective and constraint functions
2. calibration (least squares) terms and constraint functions
3. a generic response functions specification.

These correspond to (a) optimization, (b) deterministic (least squares) or stochastic (Bayesian) inversion, and (c) general-purpose analyzer methods such as parameter studies, DACE, and UQ methods, respectively. Refer to responses for additional details and examples.

Related Topics

Related Keywords

6.6.2 nonlinear_constraints

Description

Nonlinear constraints are supported by many of Dakota's optimizers. For a discussion of the difference between nonlinear and linear constraints, see the linear_constraints topic page. The Constraint Considerations section of the optimization_and_calibration page may also be of interest.

Related Topics

Related Keywords

6.7 interface

Description

Keywords and Concepts relating to the interface block, which is used to connect Dakota to external analysis codes (simulations, etc.)

Related Topics

- simulation_file_management
- workflow_management
- advanced_simulation_interfaces
Related Keywords

6.7.1 simulation_file_management

Description
empty

Related Topics
Related Keywords

6.7.2 workflow_management

Description
empty

Related Topics
Related Keywords

6.7.3 advanced_simulation_interfaces

Description
empty

Related Topics

• simulation_failure
• concurrency_and_parallelism

Related Keywords

6.7.4 simulation_failure

Description
empty

Related Topics
Related Keywords

6.7.5 concurrency_and_parallelism

Description
empty
Related Topics

Related Keywords

- **processors_per_analysis**: Specify the number of processors per analysis when Dakota is run in parallel
- **analysis_scheduling**: Specify the scheduling of concurrent analyses when Dakota is run in parallel
- **master**: Specify a dedicated master partition for parallel analysis scheduling
- **peer**: Specify a peer partition for parallel analysis scheduling
- **analysis_servers**: Specify the number of analysis servers when Dakota is run in parallel
- **asynchronous**: Specify local evaluation or analysis concurrency
- **analysis_concurrency**: Limit the number of analysis drivers within an evaluation that Dakota will schedule
- **evaluation_concurrency**: Determine how many concurrent evaluations Dakota will schedule
- **local_evaluation_scheduling**: Control how local asynchronous jobs are scheduled
- **batch**: Perform evaluations in batches
- **size**: Limit the number of evaluations in a batch
- **master**: Specify a dedicated master partition for parallel evaluation scheduling
- **peer**: Specify a peer partition for parallel evaluation scheduling
- **dynamic**: Specify dynamic scheduling in a peer partition when Dakota is run in parallel.
- **static**: Specify static scheduling in a peer partition when Dakota is run in parallel.
- **evaluation_servers**: Specify the number of evaluation servers when Dakota is run in parallel
- **processors_per_evaluation**: Specify the number of processors per evaluation server when Dakota is run in parallel
- **iterator_scheduling**: Specify the scheduling of concurrent iterators when Dakota is run in parallel
- **master**: Specify a dedicated master partition for parallel iterator scheduling
- **peer**: Specify a peer partition for parallel iterator scheduling
- **iterator_servers**: Specify the number of iterator servers when Dakota is run in parallel
- **processors_per_iterator**: Specify the number of processors per iterator server when Dakota is run in parallel
- **iterator_scheduling**: Specify the scheduling of concurrent iterators when Dakota is run in parallel
- **master**: Specify a dedicated master partition for parallel iterator scheduling
- **peer**: Specify a peer partition for parallel iterator scheduling
- **iterator_servers**: Specify the number of iterator servers when Dakota is run in parallel
- **processors_per_iterator**: Specify the number of processors per iterator server when Dakota is run in parallel
- **iterator_scheduling**: Specify the scheduling of concurrent iterators when Dakota is run in parallel
6.8 methods

Description

Keywords and Concepts relating to the method block, including discussion of the different methods and algorithms available in Dakota

Related Topics

- parameter_studies
- sensitivity_analysis_and_design_of_experiments
- uncertainty_quantification
- optimization_and_calibration

Related Keywords

6.8.1 parameter_studies

Description

Parameter studies employ deterministic designs to explore the effect of parametric changes within simulation models, yielding one form of sensitivity analysis. They can help assess simulation characteristics such as smoothness, multi-modality, robustness, and nonlinearity, which affect the choice of algorithms and controls in follow-on optimization and UQ studies.

Dakota’s parameter study methods compute response data sets at a selection of points in the parameter space. These points may be specified as a vector, a list, a set of centered vectors, or a multi-dimensional grid. Capability overviews and examples of the different types of parameter studies are provided in the Users Manual [5].
With the exception of output verbosity (a setting of silent will suppress some parameter study diagnostic output), Dakota’s parameter study methods do not make use of the method independent controls. Therefore, the parameter study documentation which follows is limited to the method dependent controls for the vector, list, centered, and multidimensional parameter study methods.

Related Topics

Related Keywords

• centered\_parameter\_study : Samples variables along points moving out from a center point
• list\_parameter\_study : Samples variables as a specified values
• multidim\_parameter\_study : Samples variables on full factorial grid of study points
• partitions : Samples variables on full factorial grid of study points
• vector\_parameter\_study : Samples variables along a user-defined vector

6.8.2 sensitivity\_analysis\_and\_design\_of\_experiments

Description

empty

Related Topics

• design\_and\_analysis\_of\_computer\_experiments
• sampling

Related Keywords

6.8.3 design\_and\_analysis\_of\_computer\_experiments

Description

Design and Analysis of Computer Experiments (DACE) methods compute response data sets at a selection of points in the parameter space. Three libraries are provided for performing these studies: DDACE, FSUDace, and PSUADE. The design of experiments methods do not currently make use of any of the method independent controls.

Related Topics

Related Keywords

• dace : Design and Analysis of Computer Experiments
• fsu\_cvt : Design of Computer Experiments - Centroidal Voronoi Tessellation
• fsu\_quasi\_mc : Design of Computer Experiments - Quasi-Monte Carlo sampling
• hammersley : Use Hammersley sequences
• psuade\_moat : Morris One-at-a-Time
6.8 methods

6.8.4 sampling

Description

Sampling techniques are selected using the `sampling` method selection. This method generates sets of samples according to the probability distributions of the uncertain variables and maps them into corresponding sets of response functions, where the number of samples is specified by the `samples` integer specification. Means, standard deviations, coefficients of variation (COVs), and 95% confidence intervals are computed for the response functions. Probabilities and reliabilities may be computed for `response_levels` specifications, and response levels may be computed for either `probability_levels` or `reliability_levels` specifications (refer to the Method Commands chapter in the Dakota Reference Manual[4] for additional information).

Currently, traditional Monte Carlo (MC) and Latin hypercube sampling (LHS) are supported by Dakota and are chosen by specifying `sample_type` as `random` or `lhs`. In Monte Carlo sampling, the samples are selected randomly according to the user-specified probability distributions. Latin hypercube sampling is a stratified sampling technique for which the range of each uncertain variable is divided into $N_s$ segments of equal probability, where $N_s$ is the number of samples requested. The relative lengths of the segments are determined by the nature of the specified probability distribution (e.g., uniform has segments of equal width, normal has small segments near the mean and larger segments in the tails). For each of the uncertain variables, a sample is selected randomly from each of these equal probability segments. These $N_s$ values for each of the individual parameters are then combined in a shuffling operation to create a set of $N_s$ parameter vectors with a specified correlation structure. A feature of the resulting sample set is that every row and column in the hypercube of partitions has exactly one sample. Since the total number of samples is exactly equal to the number of partitions used for each uncertain variable, an arbitrary number of desired samples is easily accommodated (as compared to less flexible approaches in which the total number of samples is a product or exponential function of the number of intervals for each variable, i.e., many classical design of experiments methods).

Advantages of sampling-based methods include their relatively simple implementation and their independence from the scientific disciplines involved in the analysis. The main drawback of these techniques is the large number of function evaluations needed to generate converged statistics, which can render such an analysis computationally very expensive, if not intractable, for real-world engineering applications. LHS techniques, in general, require fewer samples than traditional Monte Carlo for the same accuracy in statistics, but they still can be prohibitively expensive. For further information on the method and its relationship to other sampling techniques, one is referred to the works by McKay, et al.[61], Iman and Shortencarier[54], and Helton and Davis[47]. Note that under certain separability conditions associated with the function to be sampled, Latin hypercube sampling provides a more accurate estimate of the mean value than does random sampling. That is, given an equal number of samples, the LHS estimate of the mean will have less variance than the mean value obtained through random sampling.

Related Topics

Related Keywords

-`importance_sampling`: Importance sampling
-`sampling`: Randomly samples variables according to their distributions

6.8.5 uncertainty_quantification

Description

Dakota provides a variety of methods for propagating both aleatory and epistemic uncertainty.

At a high level, uncertainty quantification (UQ) or nondeterministic analysis is the process of characterizing input uncertainties, forward propagating these uncertainties through a computational model, and performing statistical or interval assessments on the resulting responses. This process determines the effect of uncertainties and assumptions on model outputs or results. In Dakota, uncertainty quantification methods specifically focus on the forward propagation part of the process, where probabilistic or interval information on parametric inputs are mapped through...
the computational model to assess statistics or intervals on outputs. For an overview of these approaches for engineering applications, consult [43].

UQ is related to sensitivity analysis in that the common goal is to gain an understanding of how variations in the parameters affect the response functions of the engineering design problem. However, for UQ, some or all of the components of the parameter vector, are considered to be uncertain as specified by particular probability distributions (e.g., normal, exponential, extreme value), or other uncertainty structures. By assigning specific distributional structure to the inputs, distributional structure for the outputs (i.e., response statistics) can be inferred. This migrates from an analysis that is more (*qualitative*) in nature, in the case of sensitivity analysis, to an analysis that is more rigorously (*quantitative*).

UQ methods are often distinguished by their ability to propagate aleatory or epistemic input uncertainty characterizations, where aleatory uncertainties are irreducible variabilities inherent in nature and epistemic uncertainties are reducible uncertainties resulting from a lack of knowledge. Since sufficient data is generally available for aleatory uncertainties, probabilistic methods are commonly used for computing response distribution statistics based on input probability distribution specifications. Conversely, for epistemic uncertainties, any use of probability distributions is based on subjective knowledge rather than objective data, and we may alternatively explore nonprobabilistic methods based on interval specifications.

Dakota contains capabilities for performing nondeterministic analysis with both types of input uncertainty. These UQ methods have been developed by Sandia Labs, in conjunction with collaborators in academia [31], [32], [21], [82].

The aleatory UQ methods in Dakota include various sampling-based approaches (e.g., Monte Carlo and Latin Hypercube sampling), local and global reliability methods, and stochastic expansion (polynomial chaos expansions and stochastic collocation) approaches. The epistemic UQ methods include local and global interval analysis and Dempster-Shafer evidence theory. These are summarized below and then described in more depth in subsequent sections of this chapter. Dakota additionally supports mixed aleatory/epistemic UQ via interval-valued probability, second-order probability, and Dempster-Shafer theory of evidence. These involve advanced model recursions and are described in Section.

The choice of uncertainty quantification method depends on how the input uncertainty is characterized, the computational budget, and the desired output accuracy. The recommendations for UQ methods are summarized in Table and are discussed in the remainder of the section.

TODO: Put table in Doxygen if still needed

**Related Topics**

- aleatory_uncertainty_quantification_methods
- epistemic_uncertainty_quantification_methods
- variable_support

**Related Keywords**

- adaptive_sampling: (Experimental) Adaptively refine a Gaussian process surrogate
- global_interval_est: Interval analysis using global optimization methods
- global_reliability: Global reliability methods
6.8 methods

• **gpais**: Gaussian Process Adaptive Importance Sampling
• **importance_sampling**: Importance sampling
• **local_interval_est**: Interval analysis using local optimization
• **local_reliability**: Local reliability method
• **mpp_search**: Specify which MPP search option to use
• **pof_darts**: Probability-of-Failure (POF) darts is a novel method for estimating the probability of failure based on random sphere-packing.
• **rkd_darts**: Recursive k-d (RKD) Darts: Recursive Hyperplane Sampling for Numerical Integration of High-Dimensional Functions.
• **sampling**: Randomly samples variables according to their distributions

6.8.6 aleatory_uncertainty_quantification_methods

Description

Aleatory uncertainty is also known as inherent variability, irreducible uncertainty, or randomness. Aleatory uncertainty is typically characterized using probability theory.

Related Topics

• **sampling**
• **reliability_methods**
• **stochastic_expansion_methods**

Related Keywords

• **importance_sampling**: Importance sampling

6.8.7 sampling

Description

Sampling techniques are selected using the **sampling** method selection. This method generates sets of samples according to the probability distributions of the uncertain variables and maps them into corresponding sets of response functions, where the number of samples is specified by the **samples** integer specification. Means, standard deviations, coefficients of variation (COVs), and 95% confidence intervals are computed for the response functions. Probabilities and reliabilities may be computed for either **probability_levels** or **reliability_levels** specifications (refer to the Method Commands chapter in the Dakota Reference Manual[4] for additional information).

Currently, traditional Monte Carlo (MC) and Latin hypercube sampling (LHS) are supported by Dakota and are chosen by specifying **sample_type** as **random** or **lhs**. In Monte Carlo sampling, the samples are selected randomly according to the user-specified probability distributions. Latin hypercube sampling is a stratified sampling technique for which the range of each uncertain variable is divided into \( N_s \) segments of equal probability, where \( N_s \) is the number of samples requested. The relative lengths of the segments are determined by the nature of the specified probability distribution (e.g., uniform has segments of equal width, normal has small segments near the mean and larger segments in the tails). For each of the uncertain variables, a sample is selected randomly from each of these equal probability segments. These \( N_s \) values for each of the individual parameters are then combined in a shuffling operation to create a set of \( N_s \) parameter vectors with a specified correlation structure. A feature
of the resulting sample set is that \textit{every row and column in the hypercube of partitions has exactly one sample}. Since the total number of samples is exactly equal to the number of partitions used for each uncertain variable, an arbitrary number of desired samples is easily accommodated (as compared to less flexible approaches in which the total number of samples is a product or exponential function of the number of intervals for each variable, i.e., many classical design of experiments methods).

Advantages of sampling-based methods include their relatively simple implementation and their independence from the scientific disciplines involved in the analysis. The main drawback of these techniques is the large number of function evaluations needed to generate converged statistics, which can render such an analysis computationally very expensive, if not intractable, for real-world engineering applications. LHS techniques, in general, require fewer samples than traditional Monte Carlo for the same accuracy in statistics, but they still can be prohibitively expensive. For further information on the method and its relationship to other sampling techniques, one is referred to the works by McKay, et al.[61], Iman and Shortencarier[54], and Helton and Davis[47]. Note that under certain separability conditions associated with the function to be sampled, Latin hypercube sampling provides a more accurate estimate of the mean value than does random sampling. That is, given an equal number of samples, the LHS estimate of the mean will have less variance than the mean value obtained through random sampling.

Related Topics

Related Keywords

- \texttt{importance\_sampling} : Importance sampling
- \texttt{sampling} : Randomly samples variables according to their distributions

6.8.8 \texttt{reliability\_methods}

Description

Reliability methods provide an alternative approach to uncertainty quantification which can be less computationally demanding than sampling techniques. Reliability methods for uncertainty quantification are based on probabilistic approaches that compute approximate response function distribution statistics based on specified uncertain variable distributions. These response statistics include response mean, response standard deviation, and cumulative or complementary cumulative distribution functions (CDF/CCDF). These methods are often more efficient at computing statistics in the tails of the response distributions (events with low probability) than sampling based approaches since the number of samples required to resolve a low probability can be prohibitive.

The methods all answer the fundamental question: “Given a set of uncertain input variables, $X$, and a scalar response function, $g$, what is the probability that the response function is below or above a certain level, $z$?” The former can be written as $P[g(X) \leq z] = F_g(z)$ where $F_g(z)$ is the cumulative distribution function (CDF) of the uncertain response $g(X)$ over a set of response levels. The latter can be written as $P[g(X) > z]$ and defines the complementary cumulative distribution function (CCDF).

This probability calculation involves a multi-dimensional integral over an irregularly shaped domain of interest, $D$, where $g(X) < z$ as displayed in Figure \texttt{figUQ05} for the case of two variables. The reliability methods all involve the transformation of the user-specified uncertain variables, $X$, with probability density function, $p(x_1, x_2)$, which can be non-normal and correlated, to a space of independent Gaussian random variables, $u$, possessing a mean value of zero and unit variance (i.e., standard normal variables). The region of interest, $D$, is also mapped to the transformed space to yield, $D_u$, where $g(U) < z$ as shown in Figure \texttt{figUQ06}. The Nataf transformation[17], which is identical to the Rosenblatt transformation[75] in the case of independent random variables, is used in Dakota to accomplish this mapping. This transformation is performed to make the probability calculation more tractable. In the transformed space, probability contours are circular in nature as shown in Figure \texttt{figUQ06} unlike in the original uncertain variable space, Figure \texttt{figUQ05}. Also, the multi-dimensional integrals can be approximated by simple functions of a single parameter, $\beta$, called the reliability index. $\beta$ is the minimum Euclidean distance from the origin in the transformed space to the response surface. This point is also known as the most probable point (MPP) of failure. Note, however, the methodology is equally applicable for generic functions, not simply those corresponding to failure criteria; this nomenclature is due to the origin of these methods within the disciplines of structural safety and reliability. Note that
there are local and global reliability methods. The majority of the methods available are local, meaning that a local optimization formulation is used to locate one MPP. In contrast, global methods can find multiple MPPs if they exist.

Related Topics

Related Keywords

- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
- **global_reliability**: Global reliability methods
- **u_gaussian_process**: Create GP surrogate in u-space
- **x_gaussian_process**: Create GP surrogate in x-space
- **local_reliability**: Local reliability method
- **mpp_search**: Specify which MPP search option to use
- **integration**: Integration approach
- **first_order**: First-order integration scheme
- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
- **second_order**: Second-order integration scheme
- **no_approx**: Perform MPP search on original response functions (use no approximation)
- **u_taylor_mean**: Form Taylor series approximation in "u-space" at variable means
- **u_taylor_mpp**: U-space Taylor series approximation with iterative updates
- **u_two_point**: Predict MPP using Two-point Adaptive Nonlinear Approximation in "u-space"
- **x_taylor_mean**: Form Taylor series approximation in "x-space" at variable means
- **x_taylor_mpp**: X-space Taylor series approximation with iterative updates
- **x_two_point**: Predict MPP using Two-point Adaptive Nonlinear Approximation in "x-space"
- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
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- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
- **probability_refinement**: Allow refinement of probability and generalized reliability results using importance sampling
6.8.9  stochastic_expansion_methods

Description

The development of these techniques mirrors that of deterministic finite element analysis utilizing the notions of projection, orthogonality, and weak convergence[31],[32]. Rather than estimating point probabilities, they form an approximation to the functional relationship between response functions and their random inputs, which provides a more complete uncertainty representation for use in multi-code simulations. Expansion methods include polynomial chaos expansions (PCE), which employ multivariate orthogonal polynomials that are tailored to representing particular input probability distributions, and stochastic collocation (SC), which employs multivariate interpolation polynomials. For PCE, expansion coefficients may be evaluated using a spectral projection approach (based on sampling, tensor-product quadrature, Smolyak sparse grid, or cubature methods for numerical integration) or a regression approach (least squares or compressive sensing). For SC, interpolants are formed over tensor-product or sparse grids and may be local or global, value-based or gradient-enhanced, and nodal or hierarchical. In global value-based cases (Lagrange polynomials), the barycentric formulation is used[9],[57],[50] to improve numerical efficiency and stability. Both sets of methods provide analytic response moments and variance-based metrics; however, CDF/CCDF probabilities are evaluated numerically by sampling on the expansion.

Related Topics

Related Keywords

6.8.10  epistemic_uncertainty_quantification_methods

Description

Epistemic uncertainty is uncertainty due to lack of knowledge.

In Dakota, epistemic uncertainty analysis is performed using interval analysis or Dempster-Shafer theory of evidence.

Note that epistemic uncertainty can also be modeled probabilistically. It would be more accurate to call this class of method, non-probabilistic uncertainty quantification, but the name persists for historical reasons.

Related Topics

- interval_estimation
- evidence_theory

Related Keywords

- global_evidence : Evidence theory with evidence measures computed with global optimization methods
- global_interval_est : Interval analysis using global optimization methods
- local_evidence : Evidence theory with evidence measures computed with local optimization methods
- local_interval_est : Interval analysis using local optimization

6.8.11  interval_estimation

Description

In interval analysis, one assumes that nothing is known about an epistemic uncertain variable except that its value lies somewhere within an interval. In this situation, it is NOT assumed that the value has a uniform probability of
occurring within the interval. Instead, the interpretation is that any value within the interval is a possible value or a potential realization of that variable. In interval analysis, the uncertainty quantification problem is one of determining the resulting bounds on the output (defining the output interval) given interval bounds on the inputs. Again, any output response that falls within the output interval is a possible output with no frequency information assigned to it.

We have the capability to perform interval analysis using either `global_interval_est` or `local_interval_est`. In the global approach, one uses either a global optimization method or a sampling method to assess the bounds. `global_interval_est` allows the user to specify either `lhs`, which performs Latin Hypercube Sampling and takes the minimum and maximum of the samples as the bounds (no optimization is performed) or `ego`. In the case of `ego`, the efficient global optimization method is used to calculate bounds. The `ego` method is described in Section. If the problem is amenable to local optimization methods (e.g. can provide derivatives or use finite difference method to calculate derivatives), then one can use local methods to calculate these bounds. `local_interval_est` allows the user to specify either `sqp` which is sequential quadratic programming, or `nip` which is a nonlinear interior point method.

Note that when performing interval analysis, it is necessary to define interval uncertain variables as described in Section. For interval analysis, one must define only one interval per input variable, in contrast with Dempster-Shafer evidence theory, where an input can have several possible intervals. Interval analysis can be considered a special case of Dempster-Shafer evidence theory where each input is defined by one input interval with a basic probability assignment of one. In Dakota, however, the methods are separate and semantic differences exist in the output presentation. If you are performing a pure interval analysis, we recommend using either `global_interval_est` or `local_interval_est` instead of `global_evidence` or `local_evidence`, for reasons of simplicity. An example of interval estimation is found in the `dakota/share/dakota/examples/users/cantilever_uq_global_interval.in`, and also in Section.

Note that we have kept separate implementations of interval analysis and Dempster-Shafer evidence theory because our users often want to couple interval analysis on an outer loop” with an aleatory, probabilistic analysis on an inner loop” for nested, second-order probability calculations. See Section for additional details on these nested approaches. These interval methods can also be used as the outer loop within an interval-valued probability analysis for propagating mixed aleatory and epistemic uncertainty – refer to Section for additional details.

Interval analysis is often used to model epistemic uncertainty. In interval analysis, the uncertainty quantification problem is one of determining the resulting bounds on the output (defining the output interval) given interval bounds on the inputs.

We can do interval analysis using either `global_interval_est` or `local_interval_est`. In the global approach, one uses either a global optimization method or a sampling method to assess the bounds, whereas the local method uses gradient information in a derivative-based optimization approach.

An example of interval estimation is shown in Figure, with example results in Figure. This example is a demonstration of calculating interval bounds for three outputs of the cantilever beam problem. The cantilever beam problem is described in detail in Section. Given input intervals of [1,10] on beam width and beam thickness, we can see that the interval estimate of beam weight is approximately [1,100].

```plaintext
Min and Max estimated values for each response function:
weight: Min = 1.0000169352e+00 Max = 9.9999491948e+01
stress: Min = -9.7749994284e-01 Max = 2.1499428450e+01
displ: Min = -9.9315672724e-01 Max = 6.7429714485e+01
```

---

Related Topics

- `global_interval_est` : Interval analysis using global optimization methods
- `local_interval_est` : Interval analysis using local optimization

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6.8.12 evidence_theory

Description

This section discusses Dempster-Shafer evidence theory. In this approach, one does not assign a probability distribution to each uncertain input variable. Rather, one divides each uncertain input variable into one or more intervals. The input parameters are only known to occur within intervals: nothing more is assumed.

Each interval is defined by its upper and lower bounds, and a Basic Probability Assignment (BPA) associated with that interval. The BPA represents a probability of that uncertain variable being located within that interval.

The intervals and BPAs are used to construct uncertainty measures on the outputs called "belief" and "plausibility." Belief represents the smallest possible probability that is consistent with the evidence, while plausibility represents the largest possible probability that is consistent with the evidence. For more information about the Dempster-Shafer theory of evidence, see [69] and [48].

Similar to the interval approaches, one may use global or local methods to determine plausibility and belief measures for the outputs.

Usage Notes

Note that to calculate the plausibility and belief cumulative distribution functions, one has to look at all combinations of intervals for the uncertain variables. Within each interval cell combination, the minimum and maximum value of the objective function determine the belief and plausibility, respectively. In terms of implementation, global methods use LHS sampling or global optimization to calculate the minimum and maximum values of the objective function within each interval cell, while local methods use gradient-based optimization methods to calculate these minima and maxima.

Finally, note that many non-deterministic keywords apply to the evidence methods, but one needs to be careful about the interpretation and translate probabilistic measures to epistemic ones. For example, if the user specifies distribution of type complementary, a complementary plausibility and belief function will be generated for the evidence methods (as opposed to a complementary distribution function in the sampling case). If the user specifies a set of responses levels, both the belief and plausibility will be calculated for each response level. Likewise, if the user specifies a probability level, the probability level will be interpreted both as a belief and plausibility, and response levels corresponding to the belief and plausibility levels will be calculated. Finally, if generalized reliability levels are specified, either as inputs (gen_reliability_levels) or outputs (response_levels with compute gen_reliabilities), then these are directly converted to/from probability levels and the same probability-based mappings described above are performed.

Related Topics

Related Keywords

- global_evidence : Evidence theory with evidence measures computed with global optimization methods
- local_evidence : Evidence theory with evidence measures computed with local optimization methods

6.8.13 variable_support

Description

Different nondeterministic methods have differing support for uncertain variable distributions. Tables 5.37, 5.38, and 5.39 summarize the uncertain variables that are available for use by the different methods, where a "-" indicates that the distribution is not supported by the method, a "U" means the uncertain input variables of this type must be uncorrelated, a "C" denotes that correlations are supported involving uncertain input variables of this type, and an "A" means the appropriate variables must be specified as active in the variables specification block. For example, if one wants to support sampling or a stochastic expansion method over both continuous uncertain and continuous state variables, the specification active all must be listed in the variables specification block. Additional notes include:
• we have four variants for stochastic expansions (SE), listed as Wiener, Askey, Extended, and Piecewise which draw from different sets of basis polynomials. The term stochastic expansion indicates polynomial chaos and stochastic collocation collectively, although the Piecewise option is only currently supported for stochastic collocation. Refer to polynomial_chaos and stoch_collocation for additional information on these three options.

• methods supporting the epistemic interval distributions have differing approaches: sampling and the lhs option of global_interval_est model the interval basic probability assignments (BPAs) as continuous histogram bin distributions for purposes of generating samples; local_interval_est and the ego option of global_interval_est ignore the BPA details and models these variables as simple bounded regions defined by the cell extremes; and local_evidence and global_evidence model the interval specifications as true BPAs.

<table>
<thead>
<tr>
<th>Distribution Type</th>
<th>Sampling</th>
<th>Local Reliability</th>
<th>Global Reliability</th>
<th>Wiener SE</th>
<th>Askey SE</th>
<th>Extended SE</th>
<th>Piecewise SE</th>
<th>Local Interval</th>
<th>Global Interval</th>
<th>Local Evidence</th>
<th>Global Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
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<td>-</td>
<td>-</td>
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<td>-</td>
</tr>
<tr>
<td>Bounded Normal</td>
<td>C</td>
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<td>U</td>
<td>-</td>
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<td>-</td>
</tr>
<tr>
<td>Lognormal</td>
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<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>-</td>
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</tr>
<tr>
<td>Bounded Lognormal</td>
<td>C</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Uniform</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>U</td>
<td>U</td>
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<td>-</td>
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<tr>
<td>Loguniform</td>
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<tr>
<td>Triangular</td>
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<td>U</td>
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<td>-</td>
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<tr>
<td>Exponential</td>
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<td>C</td>
<td>U</td>
<td>U</td>
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<td>-</td>
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<td>Gamma</td>
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<td>-</td>
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<td>-</td>
</tr>
<tr>
<td>Gumbel</td>
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<td>U</td>
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<td>-</td>
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</tr>
<tr>
<td>Frechet</td>
<td>C</td>
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<td>C</td>
<td>C</td>
<td>C</td>
<td>U</td>
<td>-</td>
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<td>-</td>
</tr>
<tr>
<td>Weibull</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>C</td>
<td>U</td>
<td>-</td>
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<td>-</td>
</tr>
<tr>
<td>Continuous Histogram Bin</td>
<td>C</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>U</td>
<td>-</td>
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</tr>
</tbody>
</table>

Table 6.1: Summary of Distribution Types supported by Nondeterministic Methods, Part I (Continuous Aleatory Types)
### Related Topics

**Related Keywords**

### 6.8.14 optimization_and_calibration

**Description**

Optimization algorithms work to minimize (or maximize) an objective function, typically calculated by the user simulation code, subject to constraints on design variables and responses. Available approaches in Dakota include well-tested, proven gradient-based, derivative-free local, and global methods for use in science and engineering design applications. Dakota also offers more advanced algorithms, e.g., to manage multi-objective optimization or perform surrogate-based minimization. This chapter summarizes optimization problem formulation, standard algorithms available in Dakota (mostly through included third-party libraries, see Section 6.5 of[5]), some advanced capabilities, and offers usage guidelines.

### 6.8.14.1 Optimization Formulations

This section provides a basic introduction to the mathematical formulation of optimization, problems. The primary goal of this section is to introduce terms relating to these topics, and is not intended to be a description of theory or numerical algorithms. For further details, consult[7],[34],[42],[67], and [87].

A general optimization problem is formulated as follows:
<table>
<thead>
<tr>
<th>Distribution Type</th>
<th>Sampling</th>
<th>Local Reliability</th>
<th>Global Reliability</th>
<th>Wiener SE</th>
<th>Askey SE</th>
<th>Extended SE</th>
<th>Piecewise SE</th>
<th>Local Interval</th>
<th>Global Interval</th>
<th>Local Evidence</th>
<th>Global Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Continuous Design</td>
<td>U,A</td>
<td>-</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Discrete Design</td>
<td>U,A</td>
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<td>-</td>
<td>-</td>
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<td>-</td>
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<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Design, Int Set, Real Set</td>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>Continuous State</td>
<td>U,A</td>
<td>-</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>U,A</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Discrete State</td>
<td>U,A</td>
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<tr>
<td>Design, Int Set, Real Set</td>
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</tr>
</tbody>
</table>

Table 6.3: Summary of Distribution Types supported by Nondeterministic Methods, Part III (Epistemic, Design, and State Types)

\[
\begin{align*}
\text{minimize:} & \quad f(x) \\
\text{subject to:} & \quad g_L \leq g(x) \leq g_U \\
& \quad h(x) = h_t \\
& \quad a_L \leq A_i x \leq a_U \\
& \quad A_e x = a_t \\
& \quad x_L \leq x \leq x_U
\end{align*}
\]

where vector and matrix terms are marked in bold typeface. In this formulation, \( x = [x_1, x_2, \ldots, x_n] \) is an \( n \)-dimensional vector of real-valued design variables or design parameters. The \( n \)-dimensional vectors, \( x_L \) and \( x_U \), are the lower and upper bounds, respectively, on the design parameters. These bounds define the allowable values for the elements of \( x \), and the set of all allowable values is termed the design space or the parameter space. A design point or a sample point is a particular set of values within the parameter space.

The optimization goal is to minimize the objective function, \( f(x) \), while satisfying the constraints. Constraints can be categorized as either linear or nonlinear and as either inequality or equality. The nonlinear inequality constraints, \( g(x) \), are “2-sided,” in that they have both lower and upper bounds, \( g_L \) and \( g_U \), respectively. The nonlinear equality constraints, \( h(x) \), have target values specified by \( h_t \). The linear inequality constraints create a linear system \( A_i x \), where \( A_i \) is the coefficient matrix for the linear system. These constraints are also 2-sided as they have lower and upper bounds, \( a_L \) and \( a_U \), respectively. The linear equality constraints create a linear system \( A_e x \), where \( A_e \) is the
Many different methods exist to solve the optimization problem given in Section 6.1 of[5], all of which iterate on \( x \) in some manner. That is, an initial value for each parameter in \( x \) is chosen, the response quantities, \( f(x), g(x), h(x), \) are computed, often by running a simulation, and some algorithm is applied to generate a new \( x \) that will either reduce the objective function, reduce the amount of infeasibility, or both. To facilitate a general presentation of these methods, three criteria will be used in the following discussion to differentiate them: optimization problem type, search goal, and search method.

The optimization problem type can be characterized both by the types of constraints present in the problem and by the linearity or nonlinearity of the objective and constraint functions. For constraint categorization, a hierarchy of complexity exists for optimization algorithms, ranging from simple bound constraints, through linear constraints, to full nonlinear constraints. By the nature of this increasing complexity, optimization problem categorizations are inclusive of all constraint types up to a particular level of complexity. That is, an unconstrained problem has no constraints, a bound-constrained problem has only lower and upper bounds on the design parameters, a linearly-constrained problem has both linear and bound constraints, and a nonlinearly-constrained problem may contain the full range of nonlinear, linear, and bound constraints. If all of the linear and nonlinear constraints are equality constraints, then this is referred to as an equality-constrained problem, and if all of the nonlinear constraints are inequality constraints, then this is referred to as an inequality-constrained problem. Further categorizations can be made based on the linearity of the objective and constraint functions. A problem where the objective function and all constraints are linear is called a linear programming (LP) problem. These types of problems commonly arise in scheduling, logistics, and resource allocation applications. Likewise, a problem where at least some of the objective and constraint functions are nonlinear is called a nonlinear programming (NLP) problem. These NLP problems predominate in engineering applications and are the primary focus of Dakota.

The search goal refers to the ultimate objective of the optimization algorithm, i.e., either global or local optimization. In global optimization, the goal is to find the design point that gives the lowest feasible objective function value over the entire parameter space. In contrast, in local optimization, the goal is to find a design point that is lowest relative to a “nearby” region of the parameter space. In almost all cases, global optimization will be more computationally expensive than local optimization. Thus, the user must choose an optimization algorithm with an appropriate search scope that best fits the problem goals and the computational budget.

The search method refers to the approach taken in the optimization algorithm to locate a new design point that has a lower objective function or is more feasible than the current design point. The search method can be classified as either gradient-based or nongradient-based. In a gradient-based algorithm, gradients of the response functions are computed to find the direction of improvement. Gradient-based optimization is the search method that underlies many efficient local optimization methods. However, a drawback to this approach is that gradients can be computationally expensive, inaccurate, or even nonexistent. In such situations, nongradient-based search methods may be useful. There are numerous approaches to nongradient-based optimization. Some of the more well known of these include pattern search methods (nongradient-based local techniques) and genetic algorithms (nongradient-based global techniques).

Because of the computational cost of running simulation models, surrogate-based optimization (SBO) methods are often used to reduce the number of actual simulation runs. In SBO, a surrogate or approximate model is constructed based on a limited number of simulation runs. The optimization is then performed on the surrogate model. Dakota has an extensive framework for managing a variety of local, multipoint, global, and hierarchical surrogates for use in optimization. Finally, sometimes there are multiple objectives that one may want to optimize simultaneously instead of a single scalar objective. In this case, one may employ multi-objective methods that are described in Section 6.3.1 of[5].

This overview of optimization approaches underscores that no single optimization method or algorithm works best for all types of optimization problems. Section 6.4 of[5] offers guidelines for choosing a Dakota optimization algorithm best matched to your specific optimization problem.

6.8.14.1.1 Constraint Considerations

Dakota’s input commands permit the user to specify two-sided nonlinear inequality constraints of the form \( g_{t_i} \leq g_{t_f} \), as well as nonlinear equality constraints of the form \( h_j(x) = h_j \). Some optimizers (e.g., npsol_, optpp_, soga, and moga methods) can handle these constraint forms directly, whereas other optimizers (e.g., asynch_pattern_search, dot_, and conmin_, mesh_adaptive_search) require Dakota to perform
Gradient-based methods are best suited for efficient navigation to a local minimum in the vicinity of the initial point. They are not intended to find global optima in nonconvex design spaces. Gradient-based optimization methods, see Section 6.2.3 of [5]. Gradient-based optimization methods are highly efficient, with the best convergence rates of all the local optimization methods, and are the methods of choice when the problem is smooth, unimodal, and well-behaved. However, these methods can be among the least robust when a problem exhibits nonsmooth, discontinuous, or multimodal behavior. The derivative-free methods described in Section 6.2.2 of [5] are more appropriate for problems with these characteristics.

Gradient accuracy is a critical factor for gradient-based optimizers, as inaccurate derivatives will often lead to failures in the search or pre-mature termination of the method. Analytic gradients and Hessians are ideal but often unavailable. If analytic gradient and Hessian information can be provided by an application code, a full Newton method will achieve quadratic convergence rates near the solution. If only gradient information is available and the Hessian information is approximated from an accumulation of gradient data, the superlinear convergence rates can be obtained. It is most often the case for engineering applications, however, that a finite difference method will be used by the optimization algorithm to estimate gradient values. Dakota allows the user to select the step size for these calculations, as well as choose between forward-difference and central-difference algorithms. The finite difference step size should be selected as small as possible, to allow for local accuracy and convergence, but not so small that the steps are "in the noise." This requires an assessment of the local smoothness of the response functions using, for example, a parameter study method. Central differencing will generally produce more reliable gradients than forward differencing but at roughly twice the expense.

Gradient-based methods for nonlinear optimization problems can be described as iterative processes in which a sequence of subproblems, usually which involve an approximation to the full nonlinear problem, are solved until the solution converges to a local optimum of the full problem. The optimization methods available in Dakota fall into several categories, each of which is characterized by the nature of the subproblems solved at each iteration.

### Related Topics
- local_optimization_methods
- global_optimization_methods
Related Topics

- bayesian_calibration
- nonlinear_least_squares
- advanced_optimization

Related Keywords

- dl_solver : (Experimental) Dynamically-loaded solver

6.8.15 local_optimization_methods

Description

empty

Related Topics

- unconstrained
- constrained
- sequential_quadratic_programming

Related Keywords

- coliny_cobyla : Constrained Optimization BY Linear Approximations (COBYLA)
- nlpol_sqp : NLPOL Sequential Quadratic Program
- nonlinear_cg : (Experimental) nonlinear conjugate gradient optimization
- npsol_sqp : NPSOL Sequential Quadratic Program
- optpp_cg : A conjugate gradient optimization method
- optpp_fd_newton : Finite Difference Newton optimization method
- optpp_g_newton : Newton method based least-squares calibration
- optpp_newton : Newton method based optimization
- optpp_q_newton : Quasi-Newton optimization method
- rol : Rapid Optimization Library (ROL) is a large-scale optimization package within Trilinos.

6.8.15.1 unconstrained

Description

empty
6.8 methods

Related Topics

Related Keywords

6.8.15.2 constrained

Description

empty

Related Topics

Related Keywords

• coliny_cobyla : Constrained Optimization BY Linear Approximations (COBYLA)

6.8.15.3 sequential_quadratic_programming

Description

Sequential Quadratic Programming (SQP) algorithms are a class of mathematical programming problems used to solve nonlinear optimization problems with nonlinera constraints. These methods are a generalization of Newton's method: each iteration involves minimizing a quadratic model of the problem. These subproblems are formulated as minimizing a quadratic approximation of the Lagrangian subject to linearized constraints. Only gradient information is required; Hessians are approximated by low-rank updates defined by the step taken at each iteration. It is important to note that while the solution found by an SQP method will respect the constraints, the intermediate iterates may not. SQP methods available in Dakota are dot_sqp, nlpql_sqp, nlssol_sqp, and npsol_sqp. The particular implementation in nlpql_sqp uses a variant with distributed and non-monotone line search. Thus, this variant is designed to be more robust in the presence of inaccurate or noisy gradients common in many engineering applications.

Related Topics

Related Keywords

• nlpql_sqp : NLPQL Sequential Quadratic Program
• nlssol_sqp : Sequential Quadratic Program for nonlinear least squares
• npsol_sqp : NPSOL Sequential Quadratic Program

6.8.16 global_optimization_methods

Description

empty

Related Topics

Related Keywords

• asynch_pattern_search : Pattern search, derivative free optimization method
• coliny_direct : Dividing RECTangles method
• coliny_ea : Evolutionary Algorithm
• coliny_pattern_search : Pattern search, derivative free optimization method
• efficient_global : Global Surrogate Based Optimization, a.k.a. EGO
• ncsu_direct : Dlviding RECTangles method
• soga : Single-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)

6.8.17 bayesian_calibration

Description
See the discussion of Bayesian Calibration in the Dakota User's Manual [5].

Related Topics

Related Keywords

• bayes_calibration : Bayesian calibration
• dream : DREAM (DiffeRential Evolution Adaptive Metropolis)
• chains : Number of chains in DREAM
• crossover_chain_pairs : Number of chains used in crossover.
• gr_threshold : Convergence tolerance for the Gelman-Rubin statistic
• jump_step : Number of generations a long jump step is taken
• num_cr : Number of candidate points for each crossover.
• gpmsa : (Experimental) Gaussian Process Models for Simulation Analysis (GPMSA) Bayesian calibration
• adaptive_metropolis : Use the Adaptive Metropolis MCMC algorithm
• delayed_rejection : Use the Delayed Rejection MCMC algorithm
• dram : Use the DRAM MCMC algorithm
• metropolis_hastings : Use the Metropolis-Hastings MCMC algorithm
• proposal_covariance : Defines the technique used to generate the MCMC proposal covariance.
• derivatives : Use derivatives to inform the MCMC proposal covariance.
• prior : Uses the covariance of the prior distributions to define the MCMC proposal covariance.
• muq : Markov Chain Monte Carlo algorithms from the MUQ package
• adaptive_metropolis : Use the Adaptive Metropolis MCMC algorithm
• delayed_rejection : Use the Delayed Rejection MCMC algorithm
• dram : Use the DRAM MCMC algorithm
• metropolis_hastings : Use the Metropolis-Hastings MCMC algorithm
• proposal_covariance : Defines the technique used to generate the MCMC proposal covariance.
• derivatives : Use derivatives to inform the MCMC proposal covariance.
• prior : Uses the covariance of the prior distributions to define the MCMC proposal covariance.
6.8 methods

- **queso**: Markov Chain Monte Carlo algorithms from the QUESO package
- **adaptive_metropolis**: Use the Adaptive Metropolis MCMC algorithm
- **delayed_rejection**: Use the Delayed Rejection MCMC algorithm
- **dram**: Use the DRAM MCMC algorithm
- **metropolis_hastings**: Use the Metropolis-Hastings MCMC algorithm
- **multilevel**: Use the multilevel MCMC algorithm.
- **proposal_covariance**: Defines the technique used to generate the MCMC proposal covariance.
- **derivatives**: Use derivatives to inform the MCMC proposal covariance.
- **prior**: Uses the covariance of the prior distributions to define the MCMC proposal covariance.

6.8.18 nonlinear_least_squares

Description

Dakota's least squares branch currently contains three methods for solving nonlinear least squares problems:

- **NL2SOL**, a trust-region method that adaptively chooses between two Hessian approximations (Gauss-Newton and Gauss-Newton plus a quasi-Newton approximation to the rest of the Hessian)
- **NLSSOL**, a sequential quadratic programming (SQP) approach that is from the same algorithm family as NPSOL
- **Gauss-Newton**, which supplies the Gauss-Newton Hessian approximation to the full-Newton optimizers from OPT++.

The important difference of these algorithms from general-purpose optimization methods is that the response set is defined by calibration terms (e.g. separate terms for each residual), rather than an objective function. Thus, a finer granularity of data is used by least squares solvers as compared to that used by optimizers. This allows the exploitation of the special structure provided by a sum of squares objective function.

Related Topics

Related Keywords

- **nl2sol**: Trust-region method for nonlinear least squares
- **nlssol_sqp**: Sequential Quadratic Program for nonlinear least squares

6.8.19 advanced_optimization

Description

empty

Related Topics

- **scaling**
- **multiobjective_methods**
- **surrogate_based_optimization_methods**
Related Keywords

6.8.19.1 scaling

Description
empty

Related Topics
Related Keywords

6.8.19.2 multiobjective_methods

Description
empty

Related Topics
Related Keywords

6.8.19.3 surrogate_based_optimization_methods

Description
empty

Related Topics
Related Keywords

- efficient_global : Global Surrogate Based Optimization, a.k.a. EGO
- surrogate_based_global : Global Surrogate Based Optimization
- surrogate_based_local : Local Surrogate Based Optimization

6.9 advanced_topics

Description

Advanced Dakota capabilities

Related Topics

- advanced_strategies
- advanced_model_recursion
- advanced_simulation_interfaces
- advanced_optimization
Related Keywords

6.9.1 advanced_strategies

Description
empty

Related Topics

Related Keywords

6.9.2 advanced_model_recursion

Description
empty

Related Topics

- hybrid_and_recursions_logic

Related Keywords

6.9.3 hybrid_and_recursions_logic

Description
empty

Related Topics

Related Keywords

6.9.4 advanced_simulation_interfaces

Description
empty

Related Topics

- simulation_failure
- concurrency_and_parallelism

Related Keywords

6.9.5 simulation_failure
6.9.6 concurrency_and_parallelism

Description
empty

Related Topics

Related Keywords

• processors_per_analysis : Specify the number of processors per analysis when Dakota is run in parallel
• analysis_scheduling : Specify the scheduling of concurrent analyses when Dakota is run in parallel
• master : Specify a dedicated master partition for parallel analysis scheduling
• peer : Specify a peer partition for parallel analysis scheduling
• analysis_servers : Specify the number of analysis servers when Dakota is run in parallel
• asynchronous : Specify local evaluation or analysis concurrency
• analysis_concurrency : Limit the number of analysis drivers within an evaluation that Dakota will schedule
• evaluation_concurrency : Determine how many concurrent evaluations Dakota will schedule
• local_evaluation_scheduling : Control how local asynchronous jobs are scheduled
• batch : Perform evaluations in batches
• size : Limit the number of evaluations in a batch
• master : Specify a dedicated master partition for parallel evaluation scheduling
• peer : Specify a peer partition for parallel evaluation scheduling
• dynamic : Specify dynamic scheduling in a peer partition when Dakota is run in parallel.
• static : Specify static scheduling in a peer partition when Dakota is run in parallel.
• evaluation_servers : Specify the number of evaluation servers when Dakota is run in parallel
• processors_per_evaluation : Specify the number of processors per evaluation server when Dakota is run in parallel
• iterator_scheduling : Specify the scheduling of concurrent iterators when Dakota is run in parallel
• master : Specify a dedicated master partition for parallel iterator scheduling
• peer : Specify a peer partition for parallel iterator scheduling
• iterator_servers : Specify the number of iterator servers when Dakota is run in parallel
• processors_per_iterator : Specify the number of processors per iterator server when Dakota is run in parallel
• iterator_scheduling: Specify the scheduling of concurrent iterators when Dakota is run in parallel

• master: Specify a dedicated master partition for parallel iterator scheduling

• peer: Specify a peer partition for parallel iterator scheduling

• iterator_servers: Specify the number of iterator servers when Dakota is run in parallel

• processors_per_iterator: Specify the number of processors per iterator server when Dakota is run in parallel

• iterator_scheduling: Specify the scheduling of concurrent iterators when Dakota is run in parallel

• master: Specify a dedicated master partition for parallel iterator scheduling

• peer: Specify a peer partition for parallel iterator scheduling

• iterator_servers: Specify the number of iterator servers when Dakota is run in parallel

• processors_per_iterator: Specify the number of processors per iterator server when Dakota is run in parallel

• iterator_scheduling: Specify the scheduling of concurrent iterators when Dakota is run in parallel

• master: Specify a dedicated master partition for parallel iterator scheduling

• peer: Specify a peer partition for parallel iterator scheduling

• iterator_servers: Specify the number of iterator servers when Dakota is run in parallel

• processors_per_iterator: Specify the number of processors per iterator server when Dakota is run in parallel

6.9.7 advanced_optimization

Description

empty

Related Topics

• scaling

• multiobjective_methods

• surrogate_based_optimization_methods
Related Keywords

6.9.7.1 scaling

Description
empty

Related Topics

Related Keywords

6.9.7.2 multiobjective_methods

Description
empty

Related Topics

Related Keywords

6.9.7.3 surrogate_based_optimization_methods

Description
empty

Related Topics

Related Keywords

- efficient_global : Global Surrogate Based Optimization, a.k.a. EGO
- surrogate_based_global : Global Surrogate Based Optimization
- surrogate_based_local : Local Surrogate Based Optimization

6.10 packages

Description

This topic organizes information about the different software packages (libraries) that are integrated into Dakota

Related Topics

- package_coliny
- package_conmin
- package_ddace
- package_dot
6.10 packages

- package_tsudace
- package_hopspack
- package_jega
- package_nlpql
- package_npsol
- package_optpp
- package_psuade
- package_queso
- package_scolib

Related Keywords

6.10.1 package_coliny

Description

SCOLIB (formerly known as COLINY) is a collection of nongradient-based optimizers that support the Common Optimization Library INterface (COLIN). SCOLIB optimizers currently include coliny_cobyla, coliny_direct, coliny_ea, coliny_pattern_search and coliny_solis_wets. (Yes, the input spec still has "coliny" prepended to the method name.) Additional SCOLIB information is available from https://software.sandia.gov/trac/acro.

SCOLIB solvers now support bound constraints and general nonlinear constraints. Supported nonlinear constraints include both equality and two-sided inequality constraints. SCOLIB solvers do not yet support linear constraints. Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds constraint_penalty times the sum of squares of the constraint violations to the objective function. Specific exceptions to this method for handling constraint violations are noted below. (The default value of constraint_penalty is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.)

The method independent controls for max_iterations and max_function_evaluations limit the number of major iterations and the number of function evaluations that can be performed during a SCOLIB optimization, respectively. The convergence_tolerance control defines the threshold value on relative change in the objective function that indicates convergence. The output verbosity specification controls the amount of information generated by SCOLIB: the silent, quiet, and normal settings correspond to minimal reporting from SCOLIB, whereas the verbose setting corresponds to a higher level of information, and debug outputs method initialization and a variety of internal SCOLIB diagnostics. The majority of SCOLIB’s methods perform independent function evaluations that can directly take advantage of Dakota’s parallel capabilities. Only coliny_solis_wets, coliny_cobyla, and certain configurations of coliny_pattern_search are inherently serial. The parallel methods automatically utilize parallel logic when the Dakota configuration supports parallelism. Lastly, neither speculative gradients nor linear constraints are currently supported with SCOLIB.

Some SCOLIB methods exploit parallelism through the use of Dakota’s concurrent function evaluations. The nature of the algorithms, however, limits the amount of concurrency that can be exploited. The maximum amount of evaluation concurrency that can be leveraged by the various methods is as follows:

- COBYLA: one
- DIRECT: twice the number of variables
- Evolutionary Algorithms: size of the population
- Pattern Search: size of the search pattern
- Solis-Wets: one
All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

Each of the SCOLIB methods supports the `solution_target` control, which defines a convergence criterion in which the optimizer will terminate if it finds an objective function value lower than the specified target.

### Related Topics

### Related Keywords

- `coliny_beta` : (Experimental) Coliny beta solver
- `coliny_cobyla` : Constrained Optimization BY Linear Approximations (COBYLA)
- `coliny_direct` : Dividing RECTangles method
- `coliny_ea` : Evolutionary Algorithm
- `coliny_pattern_search` : Pattern search, derivative free optimization method
- `coliny_solis_wets` : Simple greedy local search method

### 6.10.2 package_conmin

#### Description

The CONMIN library[86] is a public domain library of nonlinear programming optimizers, specifically the Fletcher--Reeves conjugate gradient (`conmin_frcg`) method for unconstrained optimization, and the method of feasible directions (`conmin_mfd`) for constrained optimization. As CONMIN was a predecessor to the DOT commercial library, the algorithm controls are very similar.

#### Related Topics

#### Related Keywords

- `conmin_mfd` : CONMIN method of feasible directions

### 6.10.3 package_ddace

#### Description

The Distributed Design and Analysis of Computer Experiments (DDACE) library provides the following DACE techniques: grid sampling (`grid`), pure random sampling (`random`), orthogonal array sampling (`oas`), latin hypercube sampling (`lhs`), orthogonal array latin hypercube sampling (`oa_lhs`), Box-Behnken (`box_benhken`), and central composite design (`central_composite`).

It is worth noting that there is some overlap in sampling techniques with those available from the nondeterministic branch. The current distinction is that the nondeterministic branch methods are designed to sample within a variety of probability distributions for uncertain variables, whereas the design of experiments methods treat all variables as having uniform distributions. As such, the design of experiments methods are well-suited for performing parametric studies and for generating data sets used in building global approximations, but are not currently suited for
assessing the effect of uncertainties characterized with probability distribution. If a design of experiments over both
design/state variables (treated as uniform) and uncertain variables (with probability distributions) is desired, then
sampling can support this with active all specified in the Variables specification block.

Related Topics

Related Keywords

- dace: Design and Analysis of Computer Experiments

6.10.4 package_dot

Description

The DOT library\cite{dotlib} contains nonlinear programming optimizers, specifically the Broyden-Fletcher-Goldfarb-Shanno
\texttt{dot_bfgs} and Fletcher-Reeves conjugate gradient \texttt{dot_frcg} methods for unconstrained optimization, and the mod-
ified method of feasible directions \texttt{dot_mmfd}, sequential linear programming \texttt{dot_slp}, and sequential quadratic
programming \texttt{dot_sqp} methods for constrained optimization. To use DOT, one of these methods must be specified.

Specialized handling of linear constraints is supported with DOT; linear constraint coefficients, bounds, and targets
can be provided to DOT at start-up and tracked internally.

Method Independent Controls - Stopping Criteria

Stopping criteria are set by: max\_iterations, max\_function\_evaluations, convergence\_tolerance, and constraint\_tolerance

Note: The convergence\_tolerance criterion must be satisfied for two consecutive iterations before DOT will
terminate.

Method Independent Controls - Output

The output verbosity specification controls the amount of information generated by DOT: the silent and quiet
settings result in header information, final results, and objective function, constraint, and parameter information
on each iteration; whereas the verbose and debug settings add additional information on gradients, search
direction, one-dimensional search results, and parameter scaling factors.

Concurrency

DOT contains no parallel algorithms which can directly take advantage of concurrent evaluations. However, if
numerical\_gradients with method\_source dakota is specified, then the finite difference function eval-
uations can be performed concurrently (using any of the parallel modes described in the Users Manual\cite{dakota}). In
addition, if speculative is specified, then gradients (dakota numerical or analytic gradients) will be
computed on each line search evaluation in order to balance the load and lower the total run time in parallel opti-
mization studies.

Related Topics

Related Keywords

- dot\_sqp: DOT Sequential Quadratic Program

6.10.5 package_fsudace

Description

The Florida State University Design and Analysis of Computer Experiments (FSUDace) library provides the following
DACE techniques: quasi-Monte Carlo sampling \texttt{fsu\_quasi\_mc} based on the Halton sequence \texttt{halton} or
the Hammersley sequence (hammersley), and Centroidal Voronoi Tessellation (fsu_cvt).

Related Topics

Related Keywords

- quality_metrics : Calculate metrics to assess the quality of quasi-Monte Carlo samples
- fsu_cvt : Design of Computer Experiments - Centroidal Voronoi Tessellation
- quality_metrics : Calculate metrics to assess the quality of quasi-Monte Carlo samples
- halton : Generate samples from a Halton sequence
- fsu_quasi_mc : Design of Computer Experiments - Quasi-Monte Carlo sampling
- halton : Generate samples from a Halton sequence
- hammersley : Use Hammersley sequences
- quality_metrics : Calculate metrics to assess the quality of quasi-Monte Carlo samples

6.10.6 package_hopspack

Description

The HOPSPACK software [72] contains the asynchronous parallel pattern search (APPS) algorithm [38]. It can handle unconstrained problems as well as those with bound constraints, linear constraints, and general nonlinear constraints.

HOPSPACK is available to the public under the GNU LGPL and the source code is included with Dakota. HOPSPACK-specific software documentation is available from https://software.sandia.gov/trac/hopspack.

Related Topics

Related Keywords

- asynch_pattern_search : Pattern search, derivative free optimization method

6.10.7 package_jega

Description

The JEGA library[19] contains two global optimization methods. The first is a Multi-objective Genetic Algorithm (MOGA) which performs Pareto optimization. The second is a Single-objective Genetic Algorithm (SOGA) which performs optimization on a single objective function. Both methods support general constraints and a mixture of real and discrete variables. The JEGA library was written by John Eddy, currently a member of the technical staff in the System Readiness and Sustainment Technologies department at Sandia National Laboratories in Albuquerque. These algorithms are accessed as moga and soga within Dakota.

Related Topics

Related Keywords

- moga : Multi-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)
- soga : Single-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)
6.10 packages

6.10.8 package_nlpql

Description

The NLPQL library includes a sequential quadratic programming (SQP) optimizer, specified as Dakota’s nlpql-sqp method, for constrained optimization. The particular implementation used is NLPQLP[77], a variant with distributed and non-monotone line search. NLPQL is commercially licensed (available from the author) and not distributed with public versions of Dakota.

Related Topics

Related Keywords

• nlpql_sqp : NLPQL Sequential Quadratic Program

6.10.9 package_npsol

Description

The NPSOL library[33] contains a sequential quadratic programming (SQP) implementation (the npsol_sqp method). SQP is a nonlinear programming optimizer for constrained minimization.

Related Topics

Related Keywords

• npsol_sqp : NPSOL Sequential Quadratic Program

6.10.10 package_optpp

Description

The OPT++ library[62] contains primarily gradient-based nonlinear programming optimizers for unconstrained, bound-constrained, and nonlinearly constrained minimization: Polak-Ribiere conjugate gradient (Dakota’s optpp_cg method), quasi-Newton (Dakota’s optpp_q_newton method), finite difference Newton (Dakota’s optpp_fd_newton method), and full Newton (Dakota’s optpp_newton method).

The conjugate gradient method is strictly unconstrained, and each of the Newton-based methods are automatically bound to the appropriate OPT++ algorithm based on the user constraint specification (unconstrained, bound-constrained, or generally-constrained). In the generally-constrained case, the Newton methods use a nonlinear interior-point approach to manage the constraints. The library also contains a direct search algorithm, PDS (parallel direct search, Dakota’s optpp_pds method), which supports bound constraints.

Controls

1. max_iterations
2. max_function_evaluations
3. convergence_tolerance
4. output
5. speculative
Concurrency

OPT++’s gradient-based methods are not parallel algorithms and cannot directly take advantage of concurrent function evaluations. However, if `numerical_gradients` with `method_source dakota` is specified, a parallel Dakota configuration can utilize concurrent evaluations for the finite difference gradient computations.

Constraints

Linear constraint specifications are supported by each of the Newton methods (`optpp_newton`, `optpp_q_newton`, `optpp_fd_newton`, and `optpp_g_newton`)

- `optpp_cg` must be unconstrained
- `optpp_pds` can be, at most, bound-constrained.

Related Topics

Related Keywords

- `optpp_cg` : A conjugate gradient optimization method
- `optpp_fd_newton` : Finite Difference Newton optimization method
- `optpp_q_newton` : Newton method based least-squares calibration
- `optpp_newton` : Newton method based optimization
- `optpp_pds` : Simplex-based derivative free optimization method
- `optpp_q_newton` : Quasi-Newton optimization method

6.10.11 package_psuade

Description

The Problem Solving Environment for Uncertainty Analysis and Design Exploration (PSUADE) is a Lawrence Livermore National Laboratory tool for metamodeling, sensitivity analysis, uncertainty quantification, and optimization. Its features include non-intrusive and parallel function evaluations, sampling and analysis methods, an integrated design and analysis framework, global optimization, numerical integration, response surfaces (MARS and higher order regressions), graphical output with Pgpplot or Matlab, and fault tolerance [84].

Related Topics

Related Keywords

- `psuade_moat` : Morris One-at-a-Time

6.10.12 package_queso

Description

QUESO stands for Quantification of Uncertainty for Estimation, Simulation, and Optimization. It supports Bayesian calibration methods. It is developed at The University of Texas at Austin.
6.10 packages

Related Topics

- bayes_calibration: Bayesian calibration
- gpmsa: (Experimental) Gaussian Process Models for Simulation Analysis (GPMSA) Bayesian calibration
- queso: Markov Chain Monte Carlo algorithms from the QUESO package

6.10.13 package_scolib

Description

SCOLIB (formerly known as COLINY) is a collection of nongradient-based optimizers that support the Common Optimization Library Interface (COLIN). SCOLIB optimizers currently include coliny_cobyla, coliny_direct, coliny_ea, coliny_pattern_search and coliny_solis_wets. (Yes, the input spec still has “coliny” prepended to the method name.) Additional SCOLIB information is available from https://software.sandia.gov/trac/acro.

SCOLIB solvers now support bound constraints and general nonlinear constraints. Supported nonlinear constraints include both equality and two-sided inequality constraints. SCOLIB solvers do not yet support linear constraints. Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds constraint_penalty times the sum of squares of the constraint violations to the objective function. Specific exceptions to this method for handling constraint violations are noted below. (The default value of constraint_penalty is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.)

The method independent controls for max_iterations and max_function_evaluations limit the number of major iterations and the number of function evaluations that can be performed during a SCOLIB optimization, respectively. The convergence_tolerance control defines the threshold value on relative change in the objective function that indicates convergence. The output verbosity specification controls the amount of information generated by SCOLIB: the silent, quiet, and normal settings correspond to minimal reporting from SCOLIB, whereas the verbose setting corresponds to a higher level of information, and debug outputs method initialization and a variety of internal SCOLIB diagnostics. The majority of SCOLIB’s methods perform independent function evaluations that can directly take advantage of Dakota’s parallel capabilities. Only coliny_solis_wets, coliny_cobyla, and certain configurations of coliny_pattern_search are inherently serial. The parallel methods automatically utilize parallel logic when the Dakota configuration supports parallelism. Lastly, neither speculative gradients nor linear constraints are currently supported with SCOLIB.

Some SCOLIB methods exploit parallelism through the use of Dakota’s concurrent function evaluations. The nature of the algorithms, however, limits the amount of concurrency that can be exploited. The maximum amount of evaluation concurrency that can be leveraged by the various methods is as follows:

- COBYLA: one
- DIRECT: twice the number of variables
- Evolutionary Algorithms: size of the population
- Pattern Search: size of the search pattern
- Solis-Wets: one

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The misc_options optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.
Each of the SCOLIB methods supports the `solution_target` control, which defines a convergence criterion in which the optimizer will terminate if it finds an objective function value lower than the specified target.

**Related Topics**

**Related Keywords**

- `coliny_beta`: (Experimental) Coliny beta solver
- `coliny_cobyla`: Constrained Optimization BY Linear Approximations (COBYLA)
- `coliny_direct`: Dividing RECTangles method
- `coliny_ea`: Evolutionary Algorithm
- `coliny_pattern_search`: Pattern search, derivative free optimization method
- `coliny_solis_wets`: Simple greedy local search method
Chapter 7
Keywords Area

This page summarizes the overall input file structure, syntax, and the six types of blocks that may appear in Dakota input. Some are optional and some may appear multiple times:

- **environment** (optional; 0 or 1 may appear)
- **method** (required; 1 or more may appear)
- **model** (optional; 0 or more may appear)
- **variables** (required; 1 or more may appear)
- **interface** (required; 1 or more may appear)
- **responses** (required; 1 or more may appear)

**Introduction to Dakota Keywords**

In Dakota, the *environment* manages execution modes and I/O streams and defines the top-level iterator. Generally speaking, an iterator contains a model and a model contains a set of *variables*, an *interface*, and a set of *responses*. An iterator repeatedly operates on the model to map the variables into responses using the interface. Each of these six components (environment, method, model, variables, interface, and responses) are separate specifications in the user's input file, and as a whole, determine the study to be performed during an execution of the Dakota software.

A Dakota execution is limited to a single environment, but may involve multiple methods and multiple models. In particular, advanced iterators (i.e., meta- and component-based iterators) and advanced models (i.e., nested and surrogate models) may specialize to include recursions with additional sub-iterators and sub-models. Since each model may contain its own variables, interface, and responses, there may be multiple specifications of the method, model, variables, interface, and responses sections.

**Keyword Pages**

Every Dakota keyword has its own page in this manual. The page describes:

- Whether the keyword takes **ARGUMENTS**, and the data type Additional notes about **ARGUMENTS** can be found here: Specifying Arguments.
- Whether it has an **ALIAS**
- Which additional keywords can be specified to change its behavior
- Which of these additional keywords are required or optional
- Additional information about how to use the keyword in an input file
7.1 environment

- Keywords Area
- environment

Top-level settings for Dakota execution

Topics

This keyword is related to the topics:

- block

Specification

**Alias:** none

**Argument(s):** none

**Default:** no environment

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>tabular_data</td>
<td></td>
<td>Write a tabular results file with variable and response history</td>
</tr>
<tr>
<td>Optional</td>
<td>output_file</td>
<td></td>
<td>Base filename for output redirection</td>
</tr>
<tr>
<td>Optional</td>
<td>error_file</td>
<td></td>
<td>Base filename for error redirection</td>
</tr>
<tr>
<td>Optional</td>
<td>read_restart</td>
<td></td>
<td>Base filename for restart file read</td>
</tr>
<tr>
<td>Optional</td>
<td>write_restart</td>
<td></td>
<td>Base filename for restart file write</td>
</tr>
<tr>
<td>Optional</td>
<td>output_precision</td>
<td></td>
<td>Control the output precision</td>
</tr>
<tr>
<td>Optional</td>
<td>results_output</td>
<td></td>
<td>(Experimental) Write a summary file containing the final results</td>
</tr>
</tbody>
</table>
## 7.1 environment

<table>
<thead>
<tr>
<th>Optional</th>
<th>graphics</th>
<th>(DEPRECATED) Display plots of variables and responses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>check</td>
<td>Invoke Dakota in input check mode</td>
</tr>
<tr>
<td>Optional</td>
<td>pre_run</td>
<td>Invoke Dakota with pre-run mode active</td>
</tr>
<tr>
<td>Optional</td>
<td>run</td>
<td>Invoke Dakota with run mode active</td>
</tr>
<tr>
<td>Optional</td>
<td>post_run</td>
<td>Invoke Dakota with post-run mode active</td>
</tr>
<tr>
<td>Optional</td>
<td>top_method_pointer</td>
<td>Identify which method leads the Dakota study</td>
</tr>
</tbody>
</table>

### Description

The environment section in a Dakota input file is optional. It specifies the top-level solution environment, optionally including run modes, output controls, and identification of the primary iterative method (`top_method_pointer`). The output-related keywords address generation of tabular and results data, and precision of numerical output.

### Run Mode Defaults

Dakota run phases include `check`, `pre_run`, `run`, and `post_run`. The default behavior is to `pre_run`, `run`, and `post_run`, though any or all of these may be specified to select specific run phases. Specifying `check` will cause Dakota to exit before any selected run modes.

#### 7.1.1 tabular_data

- **Keywords Area**
  - environment
  - tabular_data

Write a tabular results file with variable and response history

### Topics

This keyword is related to the topics:

- dakota_output

### Specification

**Alias:** tabular_graphics_data

**Argument(s):** none
Default: no tabular data output

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>Tabular Data</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Format (Group 1)</td>
<td>tabular_data_file</td>
<td>File name for tabular data output</td>
</tr>
<tr>
<td></td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Specifying the `tabular_data` flag writes to a data file the same variable and response function history data plotted when using the (deprecated) `graphics` flag. Within the generated data file, the variables and response functions appear as columns and each function evaluation provides a new table row. This capability is most useful for post-processing of Dakota results with third-party graphics tools such as MatLab, Excel, Tecplot, etc.

There is no dependence between the `graphics` flag and the `tabular_data` flag; they may be used independently or concurrently.

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

See Also

These keywords may also be of interest:

- `graphics`

7.1.1.1 tabular_data_file

- Keywords Area
- environment
- `tabular_data`
- `tabular_data_file`

File name for tabular data output
7.1 environment

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: tabular_graphics_file

Argument(s): STRING

Default: dakota_tabular.dat

Description

Specifies a name to use for the tabular data file, overriding the default dakota_tabular.dat.

7.1.1.2 custom_annotated

- Keywords Area
- environment
- tabular_data
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The **annotated** format is the default for tabular export/import. To control which header row and columns are in the input/output, specify **custom_annotated**, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to **annotated** format, though **freeform** remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
        custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.1.1.2.1 header

- Keywords Area
- environment
- tabular_data
- custom_annotated
- header

Enable header row in custom-annotated tabular file.
7.1 environment

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.1.1.2.2 eval_id

• Keywords Area
• environment
• tabular_data
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.1.1.2.3 interface_id

• Keywords Area
• environment
• tabular_data
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column
Description

See description of parent `custom_annotated`

7.1.1.3 annotated

- Keywords Area
- environment
- tabular_data
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.1.1.4 freeform

- Keywords Area
- environment
- tabular_data
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
  ...
```

7.1.2 output_file

• Keywords Area
  • environment
  • output_file

Base filename for output redirection

Topics

This keyword is related to the topics:

• dakota_IO
  • command_line_options

Specification

Alias: none
Argument(s): STRING
Default: output to console, not file

Description

Specify a base filename to which Dakota output will be directed. Output will (necessarily) be redirected after the input file is parsed. This option is overridden by any command-line -output option.

Default Behavior

Output to console (screen).
7.1.3 error_file

- Keywords Area
- environment
- error_file

Base filename for error redirection

Topics

This keyword is related to the topics:

- dakota_IO
- command_line_options

Specification

Alias: none
Argument(s): STRING
Default: errors to console, not file

Description

Specify a base filename to which Dakota errors will be directed. Errors will (necessarily) be redirected after the input file is parsed. This option is overridden by any command-line -error option.

Default Behavior

Errors to console (screen).

7.1.4 read_restart

- Keywords Area
- environment
- read_restart

Base filename for restart file read

Topics

This keyword is related to the topics:

- dakota_IO
- command_line_options
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no restart read

**Child Keywords:**
7.1 environment

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>stop_restart</td>
</tr>
</tbody>
</table>

**Description**

Specify a base filename for the restart file Dakota should read. This option is overridden by any command-line -read_restart option.

**Default Behavior**

No restart file is read.

7.1.4.1 stop_restart

- Keywords Area
- environment
- read_restart
- stop_restart

Evaluation ID number at which to stop reading restart file

**Topics**

This keyword is related to the topics:

- dakota_IO

**Specification**

Alias: none

Argument(s): INTEGER

Default: read all records

**Description**

This option is overridden by any command-line -stop_restart option.

7.1.5 write_restart

- Keywords Area
- environment
- write_restart

Base filename for restart file write
Topics

This keyword is related to the topics:

- dakota_IO
- command_line_options

Specification

Alias: none
Argument(s): STRING
Default: dakota.rst

Description

Specify a base filename for the restart file Dakota should write. This option is overridden by any command-line -write_restart option.

7.1.6 output_precision

- Keywords Area
- environment
- output_precision

Control the output precision

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none
Argument(s): INTEGER
Default: 10

Description

The precision of numeric output precision can be set with output_precision, with an upper limit of 16. When not specified, most Dakota output will default to a precision of 10, though filesystem interfaces and pre-run output use higher precision for accuracy and better results reproducibility.
7.1.7 results_output

• Keywords Area
  • environment
  • results_output

(Experimental) Write a summary file containing the final results

Topics

This keyword is related to the topics:

• dakota_output

Specification

Alias: none

Argument(s): none

Default: no results output

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>results_output_file</td>
<td>results_output_file</td>
<td>The base file name of the results file</td>
</tr>
<tr>
<td>Optional</td>
<td>text</td>
<td>text</td>
<td>Write results to file in text format</td>
</tr>
<tr>
<td>Optional</td>
<td>hdf5</td>
<td>hdf5</td>
<td>Write results to file in HDF5 format</td>
</tr>
</tbody>
</table>

Description

(Experimental)

Dakota writes final results for most methods to the screen (console). This keyword enables experimental support for writing them to disk, as well. By default, they are written in a structured text format to a file named `dakota_results.txt`. Text format can be explicitly selected using the `text` sub-keyword. If Dakota was built with HDF5 enabled, the sub-keyword `hdf5` causes results to be written in HDF5 format, as well.

The name of the results file can be changed using the `results_output_file` keyword.

Text output eventually will be deprecated, and HDF5 will be enabled in all distributed builds of Dakota. The layout of the HDF5 file is described in the Dakota HDF5 Output section of this manual. The contents, organization, and format of results files are all under active development and are subject to change.

7.1.7.1 results_output_file

• Keywords Area
  • environment
  • results_output
• results_output_file

The base file name of the results file

Topics

This keyword is related to the topics:

• dakota_output

Specification

Alias: none
Argument(s): STRING
Default: dakota_results

Description

The default filename is dakota_results.txt for text output and dakota_results.h5 for HDF5. The dakota_results stem can be changed using this keyword. Do not include the extension; these will be added by Dakota.

7.1.7.2 text

• Keywords Area

• environment

• results_output

• text

Write results to file in text format

Topics

This keyword is related to the topics:

• dakota_output

Specification

Alias: none
Argument(s): none
Default: text output
Description

When this keyword is present, Dakota will right (some) final method results to disk in a custom structured text format instead of just to the console. Support for text format results output will be deprecated in a future release, and the contents and organization of the output file is subject to change.

The default name of the file is `dakota_results.txt`. This can be changed using the `results_output_file` keyword.

Examples

```
environment
  results_output
text
  results_output_file 'my_results'  # The .txt extension will be added
```

7.1.7.3 hdf5

- Keywords Area
- environment
- results_output
- hdf5

Write results to file in HDF5 format

Topics

This keyword is related to the topics:

- dakota_output

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no HDF5 output  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>model_selection</td>
<td></td>
<td>Select the models that write evaluation data to HDF5</td>
</tr>
</tbody>
</table>
Description

When this keyword is present, Dakota will write (some) final method results to disk in HDF5 format instead of just to the console. This is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

With hdf5 selected, the default name of the output file is dakota_results.h5. This can be changed using the results_output_file keyword.

Examples

```ini
environment
  results_output
    hdf5
      results_output_file ‘my_results’  # The .h5 extension will be added
```

7.1.7.3.1 model_selection

- Keywords Area
- environment
- results_output
- hdf5
- model_selection

Select the models that write evaluation data to HDF5

Topics

This keyword is related to the topics:

- dakota_output

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

### Model Evaluation Storage Selection (Group 1)

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Model Evaluation Storage Selection</td>
<td>top_method</td>
<td>Write evaluation data only for the top-level method's model to HDF5</td>
</tr>
</tbody>
</table>

| | none | Write evaluation data for no models to HDF5 |
| | all_methods | Write evaluation data to HDF5 for all models that belong directly to methods |
| | all | Write evaluation data to HDF5 for all models |

#### Description

By default, when HDF5 output is enabled, Dakota writes evaluation data only for the model that belongs to the top-level method. This keyword group is used to override the default.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

The example below will be used to explain the effect of each keyword.

#### Examples

```plaintext
environment
  results_output
    hdf5
      # model_selection
      # top_method
      # all_methods
      # all
      # none
    results_output_file 'my_results' # The .h5 extension will be added

  method_pointer 'opt'

  method
    id_method 'opt'
      optpp_q_newton
    model_pointer 'surr'

  model
    id_model 'surr'
    surrogate global gaussian_process surfpack
    dace_method_pointer 'training'

  method
    id_method 'training'
    sampling
      seed 1234
    samples 20
    model_pointer 'truth_m'

  model
```
id_model 'truth_m'
simulation

interface
id_interface 'truth'
direct
    analysis_drivers 'text_book'

variables
continuous_design 2
    descriptors 'x1' 'x2'
    lower_bounds -2.0 -2.0
    upper_bounds  2.0  2.0

responses
objective_functions 2
    descriptors 'f1' 'f2'
analytic_gradients
no_hessians

7.1.7.3.1.1 top_method

• Keywords Area
• environment
• results_output
• hdf5
• model_selection
• top_method

Write evaluation data only for the top-level method's model to HDF5

Topics
This keyword is related to the topics:

• dakota_output

Specification

Alias: none
Argument(s): none
Default: top_method

Description
When this option is selected, evaluation data only for the model that belongs to the top-level method will be written to HDF5.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.
Examples

For the example input in the parent keyword, the following model groups would be written:

- /models/recast/RECAST_surr_RECAST_1/

The recast model, in this case, is automatically generated by Dakota to handle summing the pair of objective functions. It is the model that belongs directly to the method ‘opt’, which is the top-level method. See Sources of Evaluation Data for further explanation.

The following links would be added to methods and models sources groups:

- /methods/opt/sources/RECAST_surr_RECAST_1 → /models/recast/RECAST_surr_RECAST_1/

7.1.7.3.1.2  none

- Keywords Area
- environment
- results_output
- hdf5
- model_selection
- none

Write evaluation data for no models to HDF5

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none
Argument(s): none
Default: top_method

Description

When this option is selected, no model evaluation data is written to HDF5, and the /models group itself is not created.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.
7.1.7.3.1.3  all_methods

- Keywords Area
- environment
- results_output
- hdf5
- model_selection
- all_methods

Write evaluation data to HDF5 for all models that belong directly to methods

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none
Argument(s): none
Default: top_method

Description

When this option is selection, evaluation data for all models that belong directly to methods will be written to HDF5. Models that belong to other models (e.g. wrapped models in a recast relationship) are not stored.

By default, when HDF5 output is enabled, Dakota writes evaluation data only for the model that belongs to the top-level method. This keyword group is used to override the default.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

Examples

For the example input in the parent keyword, the following model groups would be written:

- /models/recast/RECAST_surr_RECAST_1/
- /models/simulation/truth_m/

The recast model, in this case, is automatically generated by Dakota to handle summing the pair of objective functions. It is the model that belongs directly to the method 'opt'. See Sources of Evaluation Data for further explanation.

The following links would be added to methods and models sources groups:

- /methods/opt/sources/RECAST_surr_RECAST_1 → /models/recast/RECAST_surr_RECAST_1/
7.1 environment

- /methods/training/sources/truth_m → /models/simulation/truth_m/

Depending on interface_selection, the following links may also be added to model sources groups:

- /models/simulation/truth_m/sources/truth → /interfaces/truth/truth_m/

7.1.7.3.1.4 all

- Keywords Area
- environment
- results_output
- hdf5
- model_selection
- all

Write evaluation data to HDF5 for all models

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none
Argument(s): none
Default: top_method

Description

When this option is selected, evaluation for all models will be written to HDF5.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

Examples

For the example input in the parent keyword, the following model groups would be written:

- /models/recast/RECAST_surr_RECAST_1/
- /models/simulation/truth_m/
- /models/surrogate/surr/

The recast model, in this case, is automatically generated by Dakota to handle summing the pair of objective functions. See Sources of Evaluation Data for further explanation.

The following links would be added to methods and models sources groups:
Depending on *interface_selection*, the following links may also be added to model sources groups:

- `/models/surrogate/surr/sources/APPROX_INTERFACE_1 → /interfaces/APPROX_INTERFACE_1/surr/`
- `/models/simulation/truth_m/sources/truth → /interfaces/truth/truth_m/`

### 7.1.7.3.2 *interface_selection*

- **Keywords Area**
- **environment**
- **results_output**
- **hdf5**
- **interface_selection**

Select the models that write evaluation data to HDF5

**Topics**

This keyword is related to the topics:

- **dakota_output**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Description</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Interface Evaluation Storage Selection (Group 1)</td>
<td>none</td>
<td>Write evaluation data for no interfaces to HDF5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>simulation</td>
<td>Write evaluation data only for simulation interfaces to HDF5</td>
</tr>
</tbody>
</table>
Description

By default, when HDF5 output is enabled, Dakota writes evaluation data for all simulation interfaces (interfaces of type fork, system, direct, etc, but not approximation interfaces, which are constructed by models of type surrogate). This keyword group is used to override the default.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

The example below will be used to explain the effect of each keyword.

Examples

```
environment
results_output
  hdf5
    # interface_selection
    # simulation
    # all
    # none
    results_output_file 'my_results' # The .h5 extension will be added

  method_pointer 'opt'

  method
    id_method 'opt'
    optpp_q_newton
    model_pointer 'surr'

  model
    id_model 'surr'
    surrogate global gaussian_process surfpack
dace_method_pointer 'training'

  method
    id_method 'training'
    sampling
      seed 1234
      samples 20
      model_pointer 'truth_m'

  model
    id_model 'truth_m'
    simulation

  interface
    id_interface 'truth'
    direct
      analysis_drivers 'text_book'

  variables
    continuous_design 2
      descriptors 'x1' 'x2'
      lower_bounds -2.0 -2.0
      upper_bounds 2.0 2.0

  responses
    objective_functions 2
      descriptors 'f1' 'f2'
      analytic_gradients
      no_hessians
```
7.1.7.3.2.1 none

- Keywords Area
- environment
- results_output
- hdf5
- interface_selection
- none

Write evaluation data for no interfaces to HDF5

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none

Argument(s): none

Default: simulation

Description

When this option is selected, no interface evaluation data is written to HDF5, and the /interfaces group is not created.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

7.1.7.3.2.2 simulation

- Keywords Area
- environment
- results_output
- hdf5
- interface_selection
- simulation

Write evaluation data only for simulation interfaces to HDF5

Topics

This keyword is related to the topics:

- dakota_output
7.1 environment

Specification

Alias: none
Argument(s): none
Default: simulation

Description

When this option is selected, evaluation data only for simulation interfaces will be written to HDF5. Simulation interfaces include fork, system, direct, matlab, python, scilab, and grid interfaces, but not approximation interfaces, which are automatically generated by Dakota to be used by surrogate models.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

Examples

For the example input in the parent keyword, the following interface groups would be written:

• /interfaces/truth/truth_m/

Depending on the model_selection, the following links may be added to model sources groups.

• /models/simulation/truth_m/sources/truth → /interfaces/truth/truth_m/

7.1.7.3.2.3 all

• Keywords Area
• environment
• results_output
• hdf5
• interface_selection
• all

Write evaluation data for all interfaces to HDF5

Topics

This keyword is related to the topics:

• dakota_output

Specification

Alias: none
Argument(s): none
Default: top_simulation
Description

When this option is selected, evaluation data for all interfaces, including both simulations and approximations, will be written to HDF5.

HDF5 output is an experimental feature, and the contents and organization of the output file is subject to change. The current organization and a brief explanation of HDF5 is provided in the Dakota HDF5 Output section of this manual.

Examples

For the example input in the parent keyword, the following interface groups would be written:

- /interfaces/truth/truth_m/
- /interfaces/APPROX_INTERFACE_1/surr/

Depending on the model_selection, the following links may be added to model sources groups.

- /models/simulation/truth_m/sources/truth → /interfaces/truth/truth_m/
- /models/surrogate/surr/sources/APPROX_INTERFACE_1 → /interfaces/APPROX_INTERFACE_1/surr/

7.1.8 graphics

- Keywords Area
- environment
- graphics

(DEPRECATED) Display plots of variables and responses

Topics

This keyword is related to the topics:

- dakota_output

Specification

Alias: none

Argument(s): none

Default: graphics off

Description

Activate Dakota’s legacy X Windows-based graphics feature. Dakota plotting and visualization capabilities are increasingly available in the Dakota graphical user interface (GUI); see the Dakota GUI User Manual in the documentation section of the Dakota website for additional details.

The graphics flag activates a 2D graphics window containing history plots for the variables and response functions in the study. This window is updated approximately every two seconds. Some study types such as surrogate-based optimization or local reliability specialize the use of the graphics window.
There is no dependence between the graphics flag and the tabular_data flag; they may be used independently or concurrently.

See Also

These keywords may also be of interest:

- tabular_data

7.1.9 check

- Keywords Area
- environment
- check

Invoke Dakota in input check mode

Topics

This keyword is related to the topics:

- command_line_options

Specification

Alias: none

Argument(s): none

Default: no check; proceed to run

Description

When specified, Dakota input will be parsed and the problem instantiated. Dakota will exit reporting whether any errors were found.

7.1.10 pre_run

- Keywords Area
- environment
- pre_run

Invoke Dakota with pre-run mode active

Topics

This keyword is related to the topics:

- command_line_options
Specification

Alias: none

Argument(s): none

Default: pre-run, run, post-run all executed

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>input</td>
<td></td>
<td>Base filename for pre-run mode data input</td>
</tr>
<tr>
<td>Optional</td>
<td>output</td>
<td></td>
<td>Base filename for pre-run mode data output</td>
</tr>
</tbody>
</table>

Description

When specified, Dakota execution will include the pre-run mode, which sets up methods and often generates parameter sets to evaluate. This mode is currently useful for parameter study, DACE, and Monte Carlo sampling methods.

Default Behavior

When no run modes are specified, Dakota will perform pre-run, run, and post-run phases.

7.1.10.1  input

- Keywords Area
- environment
- pre_run
- input

Base filename for pre-run mode data input

Topics

This keyword is related to the topics:

- dakota_IO

Specification

Alias: none

Argument(s): STRING

Default: no pre-run specific input read
Description

(For future expansion; not currently used by any methods.) Specify a base filename from which Dakota will read any pre-run input data. This option is overridden by any command-line -pre_run arguments.

7.1.10.2 output

- Keywords Area
  - environment
  - pre_run
  - output

Base filename for pre-run mode data output

Topics

This keyword is related to the topics:

- dakota_IO

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no pre-run specific output written

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>custom_annotated</td>
<td>freeform</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Specify a base filename to which Dakota will write any pre-run output data (typically parameter sets to be evaluated). This option is overridden by any command-line -pre_run arguments.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform
7.1.10.2.1 custom_annotated

- Keywords Area
- environment
- pre_run
- output
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips
Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.1.10.2.1.1  header

- Keywords Area
- environment
- pre_run
- output
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.1.10.2.1.2  eval_id

- Keywords Area
- environment
- pre_run
Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

Selects annotated tabular file format
7.1 environment

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.9009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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7.1.10.2.3 freeform

- Keywords Area
- environment
- pre_run
- output
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports_imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:
environment

```python
tabular_data
    tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
```

7.1.11 run

- Keywords Area
- environment
- run

Invoke Dakota with run mode active

Topics

This keyword is related to the topics:

- command_line_options

Specification

**Alias:** none

**Argument(s):** none

**Default:** pre-run, run, post-run all executed

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>input</td>
<td>Base filename for run mode data input</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>output</td>
<td>Base filename for run mode data output</td>
</tr>
</tbody>
</table>

Description

When specified, Dakota execution will include the run mode, which invokes interfaces to map parameters to responses.

**Default Behavior**

When no run modes are specified, Dakota will perform pre-run, run, and post-run phases.
7.1.11.1 input

• Keywords Area
• environment
• run
• input

Base filename for run mode data input

Topics

This keyword is related to the topics:

• dakota_IO

Specification

Alias: none
Argument(s): STRING
Default: no run specific input read

Description

(For future expansion; not currently used by any methods.) Specify a base filename from which Dakota will read any run input data, such as parameter sets to evaluate. This option is overridden by any command-line -run arguments.

7.1.11.2 output

• Keywords Area
• environment
• run
• output

Base filename for run mode data output

Topics

This keyword is related to the topics:

• dakota_IO

Specification

Alias: none
Argument(s): STRING
Default: no run specific output written
Description

(For future expansion; not currently used by any methods.) Specify a base filename to which Dakota will write any run output data (typically parameter, response pairs). This option is overridden by any command-line -run arguments.

7.1.12 post_run

- Keywords Area
- environment
- post_run

Invoke Dakota with post-run mode active

Topics

This keyword is related to the topics:

- command_line_options

Specification

Alias: none
Argument(s): none
Default: pre-run, run, post-run all executed
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>input</td>
<td>input</td>
<td>Base filename for post-run mode data input</td>
</tr>
<tr>
<td>Optional</td>
<td>output</td>
<td>output</td>
<td>Base filename for post-run mode data output</td>
</tr>
</tbody>
</table>

Description

When specified, Dakota execution will include the post-run mode, which analyzes parameter/response data sets and computes final results. This mode is currently useful for parameter study, DACE, and Monte Carlo sampling methods.

Default Behavior

When no run modes are specified, Dakota will perform pre-run, run, and post-run phases.

7.1.12.1 input

- Keywords Area
- environment
• \textit{post\_run}
• \textit{input}

Base filename for post-run mode data input

Topics

This keyword is related to the topics:

• \textit{dakota\_IO}

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: STRING

\textbf{Default}: no post-run specific input read

\textbf{Child Keywords}:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
</tr>
</tbody>
</table>

Description

Specify a base filename from which Dakota will read any post-run input data, such as parameter/response data on which to calculate final statistics. This option is overridden by any command-line \texttt{-post\_run} arguments.

\textbf{Usage Tips}

Dakota imports tabular data in one of three formats:

• \texttt{annotated} \texttt{(default)}
• \texttt{custom\_annotated}
• \texttt{freeform}

7.1.12.1 \textit{custom\_annotated}

• \textit{Keywords Area}
• \textit{environment}
• \textit{post\_run}
• \textit{input}
• \textit{custom\_annotated}

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

```
%eval_id  x1      x2      obj_fn   nln_ineq_con_1  nln_ineq_con_2
1          0.9     1.1     0.0002   0.26            0.76
2  0.90009  1.1  0.0001996404857  0.2601620081   0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081   0.760045
...
```

7.1.12.1.1.1 header

- Keywords Area
- environment
- post_run
- input
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

Alias: none

Argument(s): none

Default: no header

**Description**

See description of parent custom_annotated

7.1.12.1.1.2 eval_id

- Keywords Area
- environment
- post_run
- input
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

Alias: none

Argument(s): none

Default: no eval_id column
7.1 environment

Description

See description of parent custom_annotated

7.1.12.1.3 interface_id

- Keywords Area
- environment
- post_run
- input
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.1.12.1.2 annotated

- Keywords Area
- environment
- post_run
- input
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID  0.9 1.1  0.0002 0.26 0.76
2   NO_ID  0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.12.1.3 freeform

- Keywords Area
- environment
- post_run
- input
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
7.1 environment

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.1.12.2 output

• Keywords Area
• environment
• post_run
• output

Base filename for post-run mode data output
Topics

This keyword is related to the topics:

- dakota_IO

Specification

Alias: none

Argument(s): STRING

Default: no post-run specific output written

Description

(For future expansion; not currently used by any methods.) Specify a base filename to which Dakota will write any post-run output data. This option is overridden by any command-line -post_run arguments.

7.1.13 top_method_pointer

- Keywords Area
- environment
- top_method_pointer

Identify which method leads the Dakota study

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: method_pointer

Argument(s): STRING

Default: see discussion

Description

An optional top_method_pointer specification may be used to point to a particular method specification that will lead the Dakota analysis. The associated string must be a method identifier specified via id_method. If top_method_pointer is not used, then it will be inferred as described below (no top_method_pointer within an environment specification is treated the same as no environment specification).

Default Behavior

The top_method_pointer keyword is typically used in Dakota studies consisting of more than one method block to clearly indicate which is the leading method. This method provides the starting point for the iteration. The corresponding method specification may recurse with additional sub-method pointers in the case of "meta-iteration" (see method) or may specify a single method without recursion. Either case will ultimately result in identification
of one or more model specifications using `model_pointer`, which again may or may not involve further recursion (see `nested` and `surrogate` for recursion cases). Each of the model specifications identify the variables and responses specifications (using `variables_pointer` and `responses_pointer`) that are used to build the model, and depending on the type of model, may also identify an interface specification (for example, using `interface_pointer`). If one of these specifications does not provide an optional pointer, then that component will be constructed using the last specification parsed.

When the environment block is omitted, the top level method will be inferred as follows: When a single method is specified, there is no ambiguity and the sole method will be the top method. When multiple methods are specified, the top level method will be deduced from the hierarchical relationships implied by method pointers. If this inference is not well defined (e.g., there are multiple method specifications without any pointer relationship), then the default behavior is to employ the last method specification parsed.

**Examples**

Specify that the optimization method is the outermost method in an optimization under uncertainty study

```plaintext
environment
  top_method_pointer 'OPTIMIZATION_METHOD'
method
  id_method 'UQ_METHOD'
  ...
method
  id_method 'OPTIMIZATION_METHOD'
  ...
```

**See Also**

These keywords may also be of interest:

- `id_method`

### 7.2 method

- **Keywords Area**

- **method**

Begins Dakota method selection and behavioral settings.

**Topics**

This keyword is related to the topics:

- **block**

**Specification**

- **Alias:** none
- **Argument(s):** none
- **Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>id_method</td>
<td>Optional</td>
<td>Name the method block; helpful when there are multiple</td>
</tr>
<tr>
<td>Optional</td>
<td>output</td>
<td>Optional</td>
<td>Control how much method information is written to the screen and output file</td>
</tr>
<tr>
<td>Optional</td>
<td>final_solutions</td>
<td>Optional</td>
<td>Number of designs returned as the best solutions</td>
</tr>
<tr>
<td></td>
<td>hybrid</td>
<td>Optional</td>
<td>Strategy in which a set of methods synergistically seek an optimal design</td>
</tr>
<tr>
<td></td>
<td>multi_start</td>
<td>Required (Choose One)</td>
<td>Multi-Start Optimization Method</td>
</tr>
<tr>
<td></td>
<td>pareto_set</td>
<td>Required (Choose One)</td>
<td>Pareto set optimization</td>
</tr>
<tr>
<td></td>
<td>branch_and_bound</td>
<td>Required (Choose One)</td>
<td>(Experimental Capability) Solves a mixed integer nonlinear optimization problem</td>
</tr>
<tr>
<td></td>
<td>surrogate_based_local</td>
<td>Required (Choose One)</td>
<td>Local Surrogate Based Optimization</td>
</tr>
<tr>
<td></td>
<td>surrogate_based_global</td>
<td>Required (Choose One)</td>
<td>Global Surrogate Based Optimization</td>
</tr>
<tr>
<td></td>
<td>dot_frcg</td>
<td>Required (Choose One)</td>
<td>DOT conjugate gradient optimization method</td>
</tr>
<tr>
<td></td>
<td>dot_mmfd</td>
<td>Required (Choose One)</td>
<td>DOT modified method of feasible directions</td>
</tr>
<tr>
<td></td>
<td>dot_bfgs</td>
<td>Required (Choose One)</td>
<td>DOT BFGS optimization method</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Method (Iterative Algorithm) (Group 1)
<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dot_slp</td>
<td>DOT Sequential Linear Program</td>
</tr>
<tr>
<td>dot_sqp</td>
<td>DOT Sequential Quadratic Program</td>
</tr>
<tr>
<td>conmin_frcg</td>
<td>CONMIN conjugate gradient optimization method</td>
</tr>
<tr>
<td>conmin_mfd</td>
<td>CONMIN method of feasible directions</td>
</tr>
<tr>
<td>dl_solver</td>
<td>(Experimental) Dynamically-loaded solver</td>
</tr>
<tr>
<td>npsol_sqp</td>
<td>NPSOL Sequential Quadratic Program</td>
</tr>
<tr>
<td>nlssol_sqp</td>
<td>Sequential Quadratic Program for nonlinear least squares</td>
</tr>
<tr>
<td>nlpql_sqp</td>
<td>NLPQL Sequential Quadratic Program</td>
</tr>
<tr>
<td>optpp_cg</td>
<td>A conjugate gradient optimization method</td>
</tr>
<tr>
<td>optpp_q_newton</td>
<td>Quasi-Newton optimization method</td>
</tr>
<tr>
<td>optpp_fd_newton</td>
<td>Finite Difference Newton optimization method</td>
</tr>
<tr>
<td>optpp_g_newton</td>
<td>Newton method based least-squares calbration</td>
</tr>
<tr>
<td>optpp_newton</td>
<td>Newton method based optimization</td>
</tr>
<tr>
<td>optpp_pds</td>
<td>Simplex-based derivative free optimization method</td>
</tr>
<tr>
<td>Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>demo_tpl</td>
<td>Third-party optimization library integration demonstration.</td>
</tr>
<tr>
<td>rol</td>
<td>Rapid Optimization Library (ROL) is a large-scale optimization package within Trilinos.</td>
</tr>
<tr>
<td>asynch_pattern_search</td>
<td>Pattern search, derivative free optimization method</td>
</tr>
<tr>
<td>mesh_adaptive_search</td>
<td>Finds optimal variable values using adaptive mesh-based search</td>
</tr>
<tr>
<td>nowpac</td>
<td>Gradient-free inequality-constrained optimization using Nonlinear Optimization With Path Augmented Constraints (NOWPAC).</td>
</tr>
<tr>
<td>snowpac</td>
<td>Stochastic version of NOWPAC that incorporates error estimates and noise mitigation.</td>
</tr>
<tr>
<td>moga</td>
<td>Multi-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)</td>
</tr>
<tr>
<td>soga</td>
<td>Single-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)</td>
</tr>
<tr>
<td>coliny_pattern_search</td>
<td>Pattern search, derivative free optimization method</td>
</tr>
<tr>
<td>Algorithm</td>
<td>Description</td>
</tr>
<tr>
<td>------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>coliny_sols_wets</td>
<td>Simple greedy local search method</td>
</tr>
<tr>
<td>coliny_cobyla</td>
<td>Constrained Optimization BY Linear Approximations (COBYLA)</td>
</tr>
<tr>
<td>coliny_direct</td>
<td>Dividing RECTangles method</td>
</tr>
<tr>
<td>coliny_ea</td>
<td>Evolutionary Algorithm</td>
</tr>
<tr>
<td>coliny_beta</td>
<td>(Experimental) Coliny beta solver</td>
</tr>
<tr>
<td>nl2sol</td>
<td>Trust-region method for nonlinear least squares</td>
</tr>
<tr>
<td>nonlinear_cg</td>
<td>(Experimental) nonlinear conjugate gradient optimization</td>
</tr>
<tr>
<td>ncsu_direct</td>
<td>Dividing RECTangles method</td>
</tr>
<tr>
<td>genie_opt_darts</td>
<td>Voronoi-based high-dimensional global Lipschitzian optimization</td>
</tr>
<tr>
<td>genie_direct</td>
<td>Classical high-dimensional global Lipschitzian optimization</td>
</tr>
<tr>
<td>efficient_global</td>
<td>Global Surrogate Based Optimization, a.k.a. EGO</td>
</tr>
<tr>
<td>surrogate_based_uq</td>
<td>Generic UQ method for constructing and interrogating a surrogate model.</td>
</tr>
<tr>
<td>Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>function_train</td>
<td>UQ method leveraging a functional tensor train surrogate model.</td>
</tr>
<tr>
<td>multifidelity_function_train</td>
<td>Multifidelity uncertainty quantification using function train expansions.</td>
</tr>
<tr>
<td>multilevel_function_train</td>
<td>Multilevel uncertainty quantification using function train expansions.</td>
</tr>
<tr>
<td>polynomial_chaos</td>
<td>Uncertainty quantification using polynomial chaos expansions.</td>
</tr>
<tr>
<td>multifidelity_polynomial_chaos</td>
<td>Multifidelity uncertainty quantification using polynomial chaos expansions.</td>
</tr>
<tr>
<td>multilevel_polynomial_chaos</td>
<td>Multilevel uncertainty quantification using polynomial chaos expansions.</td>
</tr>
<tr>
<td>stoch_collocation</td>
<td>Uncertainty quantification with stochastic collocation.</td>
</tr>
<tr>
<td>multifidelity_stoch_collocation</td>
<td>Multifidelity uncertainty quantification using stochastic collocation.</td>
</tr>
<tr>
<td>sampling</td>
<td>Randomly samples variables according to their distributions.</td>
</tr>
<tr>
<td>Method</td>
<td>Description</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>multilevel_sampling</td>
<td>Multilevel sampling methods for UQ</td>
</tr>
<tr>
<td>multifidelity_sampling</td>
<td>Multifidelity sampling methods for UQ</td>
</tr>
<tr>
<td>multilevel-multifidelity_sampling</td>
<td>Multilevel-Multifidelity sampling methods for UQ</td>
</tr>
<tr>
<td>approximate_control_variate</td>
<td>Approximate control variate (ACV) sampling methods for UQ</td>
</tr>
<tr>
<td>importance_sampling</td>
<td>Importance sampling</td>
</tr>
<tr>
<td>gpais</td>
<td>Gaussian Process Adaptive Importance Sampling</td>
</tr>
<tr>
<td>adaptive_sampling</td>
<td>(Experimental) Adapтивely refine a Gaussian process surrogate</td>
</tr>
<tr>
<td>pof_darts</td>
<td>Probability-of-Failure (POF) darts is a novel method for estimating the probability of failure based on random sphere-packing.</td>
</tr>
<tr>
<td>rkd_darts</td>
<td>Recursive k-d (RKD) Darts: Recursive Hyperplane Sampling for Numerical Integration of High-Dimensional Functions.</td>
</tr>
<tr>
<td>Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>global_evidence</td>
<td>Evidence theory with evidence measures computed with global optimization methods</td>
</tr>
<tr>
<td>global_interval_est</td>
<td>Interval analysis using global optimization methods</td>
</tr>
<tr>
<td>bayes_calibration</td>
<td>Bayesian calibration</td>
</tr>
<tr>
<td>dace</td>
<td>Design and Analysis of Computer Experiments</td>
</tr>
<tr>
<td>fsu_cvrt</td>
<td>Design of Computer Experiments - Centroidal Voronoi Tessellation</td>
</tr>
<tr>
<td>psuade_moat</td>
<td>Morris One-at-a-Time</td>
</tr>
<tr>
<td>local_evidence</td>
<td>Evidence theory with evidence measures computed with local optimization methods</td>
</tr>
<tr>
<td>local_interval_est</td>
<td>Interval analysis using local optimization</td>
</tr>
<tr>
<td>local_reliability</td>
<td>Local reliability method</td>
</tr>
<tr>
<td>global_reliability</td>
<td>Global reliability methods</td>
</tr>
<tr>
<td>fsu_quasi_mc</td>
<td>Design of Computer Experiments - Quasi-Monte Carlo sampling</td>
</tr>
<tr>
<td>vector_parameter_study</td>
<td>Samples variables along a user-defined vector</td>
</tr>
<tr>
<td>list_parameter_study</td>
<td>Samples variables as a specified values</td>
</tr>
</tbody>
</table>
### Description

The **method** keyword signifies the start of a block in the Dakota input file. A method block contains the various keywords necessary to select a method and to control its behavior.

### Method Block Requirements

At least one **method** block must appear in the Dakota input file. Multiple **method** blocks may be needed to fully define advanced analysis approaches.

Each **method** block must specify one method and, optionally, any associated keywords that govern the behavior of the method.

### The Methods

Each **method** block must select one method.

Starting with Dakota v6.0, the methods are grouped into two types: standard methods and multi-component methods.

The standard methods are stand-alone and self-contained in the sense that they only require a model to perform a study. They do not call other methods. While methods such as `polynomial_chaos` and `efficient_global` internally utilize multiple iterator and surrogate model components, these components are generally hidden from user control due to restrictions on modularity; thus, these methods are stand-alone.

The multi-component group of methods provides a higher level "meta-algorithm" that points to other methods and models that support sub-iteration. For example, in a sequential hybrid method, the `hybrid` method specification must identify a list of subordinate methods, and the "meta-algorithm" executes these methods in sequence and transfers information between them. Surrogate-based minimizers provide another example in that they point both to other methods (e.g. what optimization method is used to solve the approximate subproblem) as well as to models (e.g. what type of surrogate model is employed). Multi-component methods generally provide some level of "plug and play" modularity, through their flexible support of a variety of method and model selections.

### Component-Based Iterator Commands

Component-based iterator specifications include hybrid, multi-start, pareto set, surrogate-based local, surrogate-based global, and branch and bound methods. Whereas a standard iterator specification only needs an optional model pointer string (specified with `model_pointer`), component-based iterator specifications can include method pointer, method name, and model pointer specifications in order to define the components employed in the "meta-iteration." In particular, these specifications identify one or more methods (by pointer or by name) to specify the subordinate iterators that will be used in the top-level algorithm. Identifying a sub-iterator by name instead of by pointer is a lightweight option that relaxes the need for a separate method specification for the sub-iterator; however, a model pointer may be required in this case to provide the specification connectivity normally supported by the method pointer. Refer to these individual method descriptions for specific requirements for these advanced methods.

### Method Independent Controls

In addition to the method, there are 10 optional keywords, which are referred to as method independent controls. These controls are valid for enough methods that it was reasonable to pull them out of the method dependent blocks.
and consolidate the specifications, however, they are NOT universally respected by all methods.

Examples

Several examples follow. The first example shows a minimal specification for an optimization method.

```
method
dot_sqp
```

This example uses all of the defaults for this method.

A more sophisticated example would be

```
method,
id_method = 'NLP1'
dot_sqp
max_iterations = 50
convergence_tolerance = 1e-4
output verbose
model_pointer = 'M1'
```

This example demonstrates the use of identifiers and pointers as well as some method independent and method dependent controls for the sequential quadratic programming (SQP) algorithm from the DOT library. The `max_iterations`, `convergence_tolerance`, and `output` settings are method independent controls, in that they are defined for a variety of methods.

The next example shows a specification for a least squares method.

```
method
  optpp_g_newton
  max_iterations = 10
  convergence_tolerance = 1.e-8
  search_method trust_region
  gradient_tolerance = 1.e-6
```

Some of the same method independent controls are present along with several method dependent controls (`search_method` and `gradient_tolerance`) which are only meaningful for OPT++ methods (see `package-optpp`).

The next example shows a specification for a nondeterministic method with several method dependent controls (refer to `sampling`).

```
method
  sampling
  samples = 100
  seed = 12345
  sample_type lhs
  response_levels = 1000. 500.
```

The last example shows a specification for a parameter study method where, again, each of the controls are method dependent (refer to `vector_parameter_study`).

```
method
  vector_parameter_study
  step_vector = 1. 1. 1.
  num_steps = 10
```

7.2.1 id_method

- Keywords Area
- method
- id_method

Name the method block; helpful when there are multiple
7.2 method

Topics
This keyword is related to the topics:

- block_identifier
- method_independent_controls

Specification
Alias: none
Argument(s): STRING
Default: strategy use of last method parsed

Description
The method identifier string is supplied with `id_method` and is used to provide a unique identifier string for use with environment or meta-iterator specifications (refer to `environment`). It is appropriate to omit a method identifier string if only one method is included in the input file, since the single method to use is unambiguous in this case.

7.2.2 output

- Keywords Area
- method
- output

Control how much method information is written to the screen and output file

Topics
This keyword is related to the topics:

- dakota_output
- method_independent_controls

Specification
Alias: none
Argument(s): none
Default: normal
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Output Level (Group 1)</td>
<td>debug</td>
<td>Level 5 of 5 - maximum</td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

Choose from a total of five output levels during the course of a Dakota study. If there is no user specification for output verbosity, then the default setting is normal.

Specific mappings are as follows:

- **silent** (i.e., really quiet): silent iterators, silent model, silent interface, quiet approximation, quiet file operations
- **quiet**: quiet iterators, quiet model, quiet interface, quiet approximation, quiet file operations
- **normal**: normal iterators, normal model, normal interface, quiet approximation, quiet file operations
- **verbose**: verbose iterators, normal model, verbose interface, verbose approximation, verbose file operations
- **debug** (i.e., really verbose): debug iterators, normal model, debug interface, verbose approximation, verbose file operations

Note that iterators and interfaces utilize the full granularity in verbosity, whereas models, approximations, and file operations do not. With respect to iterator verbosity, different iterators implement this control in slightly different ways (as described below in the method independent controls descriptions for each iterator), however the meaning is consistent.

For models, interfaces, approximations, and file operations, **quiet** suppresses parameter and response set reporting and **silent** further suppresses function evaluation headers and scheduling output. Similarly, **verbose** adds file management, approximation evaluation, and global approximation coefficient details, and **debug** further adds diagnostics from nonblocking schedulers.

### 7.2.2.1 debug

- **Keywords Area**
- **method**
- **output**
- **debug**

Level 5 of 5 - maximum

Specification

**Alias**: none

**Argument(s)**: none

Description

This is described on **output**
7.2.2.2 verbose

- Keywords Area
- method
- output
- verbose

Level 4 of 5 - more than normal

Specification

Alias: none
Argument(s): none

Description

This is described on output

7.2.2.3 normal

- Keywords Area
- method
- output
- normal

Level 3 of 5 - default

Specification

Alias: none
Argument(s): none

Description

This is described on output

7.2.2.4 quiet

- Keywords Area
- method
- output
- quiet

Level 2 of 5 - less than normal
Specification

Alias: none
Argument(s): none

Description

This is described on output

7.2.2.5 silent

- Keywords Area
- method
- output
- silent

Level 1 of 5 - minimum

Specification

Alias: none
Argument(s): none

Description

This is described on output

7.2.3 final_solutions

- Keywords Area
- method
- final_solutions

Number of designs returned as the best solutions

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Dakota Version 6.15 Reference Manual generated on November 5, 2021
7.2 method

Description

The `final_solutions` controls the number of final solutions returned by the iterator as the best solutions. For most optimizers, this is one, but some optimizers can produce multiple solutions (e.g. genetic algorithms). When using a hybrid strategy, the number of final solutions dictates how many solutions are passed from one method to another.

Examples

In the case of sampling methods, if one specifies 100 samples (for example) but also specifies `final_solutions = 5`, the five best solutions (in order of lowest response function value) are returned.

7.2.4 hybrid

- Keywords Area
- method
- hybrid

Strategy in which a set of methods synergistically seek an optimal design

Specification

Alias: none  
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Hybrid Method Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>sequential</td>
<td>Methods are run one at a time, in sequence</td>
</tr>
<tr>
<td></td>
<td></td>
<td>embedded</td>
<td>A subordinate local method provides periodic refinements to a top-level global method</td>
</tr>
<tr>
<td></td>
<td></td>
<td>collaborative</td>
<td>Multiple methods run concurrently and share information</td>
</tr>
</tbody>
</table>

Description

In a hybrid minimization method (hybrid), a set of methods synergistically seek an optimal design. The relationships among the methods are categorized as:

- collaborative
- embedded
- sequential
The goal in each case is to exploit the strengths of different optimization and nonlinear least squares algorithms at different stages of the minimization process. Global + local hybrids (e.g., genetic algorithms combined with nonlinear programming) are a common example in which the desire for identification of a global optimum is balanced with the need for efficient navigation to a local optimum.

7.2.4.1 sequential

- Keywords Area
- method
- hybrid
- sequential

Methods are run one at a time, in sequence

Specification

**Alias:** uncoupled

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sub-method Selection (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>method_name_list</td>
<td>method_pointer_list</td>
<td>List of Dakota methods to sequentially or collaboratively run</td>
</tr>
<tr>
<td>Optional</td>
<td>iterator_servers</td>
<td>Specify the number of iterator servers when Dakota is run in parallel</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>iterator_scheduling</td>
<td>Specify the scheduling of concurrent iterators when Dakota is run in parallel</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>processors_per_iterator</td>
<td>Specify the number of processors per iterator server when Dakota is run in parallel</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

In the **sequential** approach, methods are run one at a time, in sequence. The best solutions from one method are used to initialize the next method.

The sequence of methods (i.e., iterators) to run are specified using either a `method_pointer_list` or a `method_name_list` (with optional `model_pointer_list`). Any number of iterators may be specified.

Method switching is managed through the separate convergence controls of each method. The number of solutions transferred between methods is specified by the particular method through its `final_solutions` method control.
For example, if one sets up a two-level study with a first method that generates multiple solutions such as a genetic algorithm, followed by a second method that is initialized only at a single point such as a gradient-based algorithm, it is possible to take the multiple solutions generated by the first method and create several instances of the second method, each one with a different initial starting point.

The logic governing the transfer of multiple solutions between methods is as follows:

- If one solution is returned from method A, then one solution is transferred to method B.
- If multiple solutions are returned from method A, and method B can accept multiple solutions as input (for example, as a genetic algorithm population), then one instance of method B is initialized with multiple solutions.
- If multiple solutions are returned from method A but method B only can accept one initial starting point, then method B is run several times, each one with a separate starting point from the results of method A.

### 7.2.4.1.1 method_name_list

- **Keywords Area**
  - method
  - hybrid
  - sequential
  - method_name_list

List of Dakota methods to sequentially or collaboratively run

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer_list</td>
<td>Associate models with method names</td>
</tr>
</tbody>
</table>

**Description**

`method_name_list` specifies a list of Dakota methods (e.g. soga, conmin_frcg) that will be run by a hybrid sequential or hybrid collaborative method. The methods are executed with default options. The optional `model_pointer_list` may be used to associate a model with each method.

### 7.2.4.1.1.1 model_pointer_list

- **Keywords Area**
  - method
  - hybrid
  - sequential
  - method_name_list
  - model_pointer_list

Associate models with method names

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Keywords Area

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRINGLIST

Description

Using the optional keyword model_pointer_list, models can be assigned to methods specified in the method_name_list. Models are referred to by name (i.e. by their id_model labels). The length of the model_pointer_list must be either 1 or match the length of the method_name_list. If the former, the same model will be used for all methods, and if the latter, methods and models will be paired in the order that they appear in the two lists.

7.2.4.1.2 method_pointer_list

- Keywords Area
- method
- hybrid
- sequential
- method_pointer_list

Pointers to methods to execute sequentially or collaboratively

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRINGLIST

Description

method_pointer_list specifies by name the methods that are to be executed by a hybrid sequential or hybrid collaborative method. Its argument is a list of strings that refer to method blocks by name (i.e. to their id_method labels).
7.2.4.1.3 iterator_servers

- Keywords Area
- method
- hybrid
- sequential
- iterator_servers

Specify the number of iterator servers when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional iterator_servers specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.1.4 iterator_scheduling

- Keywords Area
- method
- hybrid
- sequential
- iterator_scheduling

Specify the scheduling of concurrent iterators when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>master</td>
<td>peer</td>
<td>Specify a peer partition for parallel iterator scheduling</td>
</tr>
</tbody>
</table>

**Description**

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `iterator_scheduling` specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to `ParallelLibrary` and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.1.4.1 master

- **Keywords Area**

- method

- hybrid

- sequential

- `iterator_scheduling`

- master

Specify a dedicated master partition for parallel iterator scheduling

**Topics**

This keyword is related to the topics:

- concurrency_and_parallelism

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.
7.2.4.1.4.2 peer

Specify a peer partition for parallel iterator scheduling

Topics
This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): none

Description
This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.

7.2.4.1.5 processors_per_iterator

Specify the number of processors per iterator server when Dakota is run in parallel

Topics
This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): INTEGER
Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `processors_per_iterator` specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.

7.2.4.2 embedded

- Keywords Area
  - method
  - hybrid
  - embedded

A subordinate local method provides periodic refinements to a top-level global method

Specification

**Alias**: coupled

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Global Sub-method Selection (Group 1)</td>
<td>global_method_-name</td>
<td>Specify the global method by Dakota name</td>
</tr>
<tr>
<td></td>
<td></td>
<td>global_method_-pointer</td>
<td>Pointer to global method</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Local Sub-method Selection (Group 2)</td>
<td>local_method_-name</td>
<td>Specify the local method by Dakota name</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_method_-pointer</td>
<td>Pointer to local method</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>local_search_-probability</td>
<td>Probability of executing local searches</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>iterator_servers</td>
<td>Specify the number of iterator servers when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>iterator_scheduling</td>
<td>Specify the scheduling of concurrent iterators when Dakota is run in parallel</td>
</tr>
</tbody>
</table>
In the embedded approach, a tightly-coupled hybrid is employed in which a subordinate local method provides periodic refinements to a top-level global method. Global and local method strings supplied with the `global_method_pointer` and `local_method_pointer` specifications identify the two methods to be used. Alternatively, Dakota method names (e.g. 'soga') can be supplied using the `global_method_name` and `local_method_name` keywords, which each have optional model pointer specifications. The `local_search_probability` setting is an optional specification for supplying the probability (between 0.0 and 1.0) of employing local search to improve estimates within the global search.

### 7.2.4.2.1 global_method_name

- **Keywords Area**
- **method**
- **hybrid**
- **embedded**
- **global_method_name**

 Specify the global method by Dakota name

**Specification**

**Alias:** none  
**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td><code>global_model_pointer</code></td>
<td>Pointer to model used by global method</td>
</tr>
</tbody>
</table>

**Description**

global_method_name is used to specify the global method in a hybrid embedded optimization by Dakota name (e.g. 'soga'). The name of the method is provided as a string. The method is executed with default options.

### 7.2.4.2.1.1 global_model_pointer

- **Keywords Area**
- **method**
- **hybrid**
- **embedded**
7.2 method

- global_method_name
- global_model_pointer

Pointer to model used by global method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Description

global_model_pointer can be used to specify a model for use with the Dakota method named by the global_method_name specification. The argument is a string that refers to the id_model label of the desired model.

7.2.4.2.2 global_method_pointer

- Keywords Area
- method
- hybrid
- embedded
- global_method_pointer

Pointer to global method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Description

The global_method_pointer identifies the method block to use as the global method in a hybrid embedded optimization using its id_method label.
7.2.4.2.3  local_method_name

- Keywords Area
- method
- hybrid
- embedded
- local_method_name

Specify the local method by Dakota name

**Specification**

**Alias:** none

**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>local_model_pointer</td>
<td>Pointer to model used by local method</td>
</tr>
</tbody>
</table>

**Description**

`local_method_name` is used to specify the local method in a hybrid embedded optimization by Dakota name (e.g. 'conmin_mfd'). The name of the method is provided as a string. The method is executed with default options.

7.2.4.2.3.1  local_model_pointer

- Keywords Area
- method
- hybrid
- embedded
- local_method_name
- local_model_pointer

Pointer to model used by local method

**Topics**

This keyword is related to the topics:

- `block_pointer`

**Specification**

**Alias:** none

**Argument(s):** STRING
Description

`local_model_pointer` can be used to specify a model for use with the Dakota method named by the `local_method_name` specification. The argument is a string that refers to the `id_model` label of the desired model.

7.2.4.2.4 `local_method_pointer`

- Keywords Area
- `method`
- `hybrid`
- `embedded`
- `local_method_pointer`

Pointer to local method

Topics

This keyword is related to the topics:

- `block_pointer`

Specification

**Alias**: none

**Argument(s)**: STRING

Description

The `local_method_pointer` identifies the method block to use as the local method in a hybrid embedded optimization using its `id_method`.

7.2.4.2.5 `local_search_probability`

- Keywords Area
- `method`
- `hybrid`
- `embedded`
- `local_search_probability`

Probability of executing local searches

Specification

**Alias**: none

**Argument(s)**: REAL
Description

The `local_search_probability` setting is an optional specification for supplying the probability (between 0.0 and 1.0) of employing local search to improve estimates within the global search. Its default value is 0.1.

7.2.4.2.6 iterator_servers

Specify the number of iterator servers when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

**Alias:** none

**Argument(s):** INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `iterator_servers` specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.2.7 iterator_scheduling

Specify the scheduling of concurrent iterators when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- concurrency_and_parallelism
7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>master</td>
<td>Specify a dedicated master partition for parallel iterator scheduling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel iterator scheduling</td>
</tr>
</tbody>
</table>

**Description**

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `iterator_scheduling` specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.2.7.1 master

- **Keywords Area**
- **method**
- **hybrid**
- **embedded**
- **iterator_scheduling**
- **master**

Specify a dedicated master partition for parallel iterator scheduling

**Topics**

This keyword is related to the topics:

- **concurrency_and_parallelism**

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.
7.2.4.2.7.2 peer

Specify a peer partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.

7.2.4.2.8 processors_per_iterator

Specify the number of processors per iterator server when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): INTEGER
7.2 Method

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `processors_per_iterator` specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterations. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.

7.2.4.3 collaborative

- Keywords Area
  - method
  - hybrid
  - collaborative

Multiple methods run concurrently and share information

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sub-method Selection (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>method_name_list</td>
<td>method_pointer_list</td>
<td>List of Dakota methods to sequentially or collaboratively run</td>
</tr>
<tr>
<td>Optional</td>
<td>iterator_servers</td>
<td></td>
<td>Specify the number of iterator servers when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>iterator_scheduling</td>
<td></td>
<td>Specify the scheduling of concurrent iterators when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>processors_per_iterator</td>
<td></td>
<td>Specify the number of processors per iterator server when Dakota is run in parallel</td>
</tr>
</tbody>
</table>

Description

In the collaborative approach, multiple methods work together and share solutions while executing concurrently. A list of method strings specifies the pool of iterators to be used. Any number of iterators may be specified. The
method collaboration logic follows that of either the Agent-Based Optimization or HOPSPACK codes and is currently under development and not available at this time.

### 7.2.4.3.1 method_name_list

- **Keywords Area**
  - method
  - hybrid
  - collaborative
  - method_name_list

List of Dakota methods to sequentially or collaboratively run

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>model_pointer_list</td>
<td>model_pointer_list</td>
<td>Associate models with method names</td>
</tr>
</tbody>
</table>

**Description**

`method_name_list` specifies a list of Dakota methods (e.g. `soga`, `conmin_frcg`) that will be run by a hybrid sequential or hybrid collaborative method. The methods are executed with default options. The optional `model_pointer_list` may be used to associate a model with each method.

### 7.2.4.3.1.1 model_pointer_list

- **Keywords Area**
  - method
  - hybrid
  - collaborative
  - method_name_list
  - model_pointer_list

Associate models with method names

**Topics**

This keyword is related to the topics:

- `block_pointer`
7.2 method

Specification

Alias: none
Argument(s): STRINGLIST

Description

Using the optional keyword model_pointer_list, models can be assigned to methods specified in the method_name_list. Models are referred to by name (i.e. by their id_model labels). The length of the model_pointer_list must be either 1 or match the length of the method_name_list. If the former, the same model will be used for all methods, and if the latter, methods and models will be paired in the order that they appear in the two lists.

7.2.4.3.2 method_pointer_list

- Keywords Area
- method
- hybrid
- collaborative
- method_pointer_list

Pointers to methods to execute sequentially or collaboratively

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none
Argument(s): STRINGLIST

Description

method_pointer_list specifies by name the methods that are to be executed by a hybrid sequential or hybrid collaborative method. Its argument is a list of strings that refer to method blocks by name (i.e. to their id_method labels).

7.2.4.3.3 iterator_servers

- Keywords Area
- method
- hybrid
- collaborative
- iterator_servers

Specify the number of iterator servers when Dakota is run in parallel.
Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional iterator_servers specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.3.4 iterator_scheduling

- Keywords Area
- method
- hybrid
- collaborative
- iterator_scheduling

Specify the scheduling of concurrent iterators when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>master</td>
<td>master</td>
<td>Specify a dedicated master partition for parallel iterator scheduling</td>
</tr>
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</table>
Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `iterator_scheduling` specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.4.3.4.1 master

- Keywords Area
- method
- hybrid
- collaborative
- iterator_scheduling
- master

Specify a dedicated master partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.

7.2.4.3.4.2 peer

- Keywords Area
- method
- hybrid
- collaborative
- iterator_scheduling
• **peer**

Specify a peer partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

• **concurrency_and_parallelism**

Specification

**Alias:** none

**Argument(s):** none

**Description**

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of `evaluation_scheduling`, it is not possible to specify `static` or `dynamic`.

### 7.2.4.3.5 processors_per_iterator

• **Keywords Area**
• **method**
• **hybrid**
• **collaborative**
• **processors_per_iterator**

Specify the number of processors per iterator server when Dakota is run in parallel

Topics

This keyword is related to the topics:

• **concurrency_and_parallelism**

Specification

**Alias:** none

**Argument(s):** INTEGER

**Description**

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `processors_per_iterator` specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.
7.2.5 multi_start

- Keywords Area
- method
- multi_start

Multi-Start Optimization Method

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tr>
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<td>Sub-method Selection (Group 1)</td>
<td>method_name</td>
<td>Specify sub-method by name</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>method_pointer</td>
<td>Pointer to sub-method to run from each starting point</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>random_starts</td>
<td>Number of random starting points</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>starting_points</td>
<td>List of user-specified starting points</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>iterator_servers</td>
<td>Specify the number of iterator servers when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>iterator_scheduling</td>
<td>Specify the scheduling of concurrent iterators when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>processors_per_iterator</td>
<td>Specify the number of processors per iterator server when Dakota is run in parallel</td>
</tr>
</tbody>
</table>

**Description**

In the multi-start iteration method (multi_start), a series of iterator runs are performed for different values of parameters in the model. A common use is for multi-start optimization (i.e., different local optimization runs from different starting points for the design variables), but the concept and the code are more general. Multi-start iteration is implemented within the MetaIterator branch of the Iterator hierarchy within the ConcurrentMetaIterator class. Additional information on the multi-start algorithm is available in the Users Manual[5].

The multi_start meta-iterator must specify a sub-iterator using either a method_pointer or a method_name plus optional model_pointer. This iterator is responsible for completing a series of iterative analyses from a set of different starting points. These starting points can be specified as follows: (1) using random_starts, for which the specified number of starting points are selected randomly within the variable bounds, (2)
using `starting_points`, in which the starting values are provided in a list, or (3) using both `random_starts` and `starting_points`, for which the combined set of points will be used. In aggregate, at least one starting point must be specified. The most common example of a multi-start algorithm is multi-start optimization, in which a series of optimizations are performed from different starting values for the design variables. This can be an effective approach for problems with multiple minima.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the starting points for each sub-iterator it runs, as well as the best parameters and responses returned by each sub-iterator. See the [Multistart and Pareto Set](https://dakota.org/doc/Multistart.html) documentation for details.

### 7.2.5.1 method_name

- **Keywords Area**
- **method**
- **multi_start**
- **method_name**

Specify sub-method by name

**Specification**

**Alias:** none

**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

The `method_name` keyword is used to specify a sub-method by Dakota method name (e.g. `npsol_sqp`) rather than block pointer. The method will be executed using its default settings. The optional `model_pointer` specification can be used to associate a model block with the method.

#### 7.2.5.1.1 model_pointer

- **Keywords Area**
- **method**
- **multi_start**
- **method_name**
- **model_pointer**

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

- `block_pointer`

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling, 
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                         0.1 0.2 0.6
                         0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
```
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system async evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.5.2 method_pointer

- Keywords Area
- method
- multi_start
- method_pointer

Pointer to sub-method to run from each starting point

Topics
This keyword is related to the topics:

- block_pointer

Specification

Alias: none
Argument(s): STRING

Description
The method_pointer keyword is used to specify a pointer to the sub-method block that will be run from each starting point.

7.2.5.3 random_starts

- Keywords Area
- method
- multi_start
- random_starts

Number of random starting points
7.2 method 281

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
</tbody>
</table>

Description

The multi_start meta-iterator must specify a sub-iterator using either a method_pointer or a method_name plus optional model_pointer. This iterator is responsible for completing a series of iterative analyses from a set of different starting points. These starting points can be specified as follows: (1) using random_starts, for which the specified number of starting points are selected randomly within the variable bounds, (2) using starting_points, in which the starting values are provided in a list, or (3) using both random_starts and starting_points, for which the combined set of points will be used.

7.2.5.3.1 seed

- Keywords Area
- method
- multi_start
- random_starts
- seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.
Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.5.4 starting_points

- Keywords Area
- method
- multi_start
- starting_points

List of user-specified starting points

Specification

Alias: none

Argument(s): REALLIST

Description

The `multi_start` meta-iterator must specify a sub-iterator using either a `method_pointer` or a `method_name` plus optional `model_pointer`. This iterator is responsible for completing a series of iterative analyses from a set of different starting points. These starting points can be specified as follows: (1) using `random_starts`, for which the specified number of starting points are selected randomly within the variable bounds, (2) using `starting_points`, in which the starting values are provided in a list, or (3) using both `random_starts` and `starting_points`, for which the combined set of points will be used.

7.2.5.5 iterator_servers

- Keywords Area
- method
- multi_start
- iterator_servers

Specify the number of iterator servers when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER
7.2 method

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional \texttt{iterator_servers} specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, \texttt{hybrid}, \texttt{multi\_start}, and \texttt{pareto\_set} component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.5.6 \texttt{iterator\_scheduling}

- \textit{Keywords Area}
- \textit{method}
- \textit{multi\_start}
- \textit{iterator\_scheduling}

Specify the scheduling of concurrent iterators when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- \textit{concurrency\_and\_parallelism}

Specification

\textbf{Alias:} none

\textbf{Argument(s):} none

\textbf{Child Keywords:}

<table>
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<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>master</td>
<td>Specify a dedicated master partition for parallel iterator scheduling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel iterator scheduling</td>
</tr>
</tbody>
</table>

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional \texttt{iterator\_scheduling} specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, \texttt{hybrid}, \texttt{multi\_start}, and \texttt{pareto\_set} component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.
7.2.5.6.1 master

- Keywords Area
- method
- multi_start
- iterator_scheduling
- master

Specify a dedicated master partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.

7.2.5.6.2 peer

- Keywords Area
- method
- multi_start
- iterator_scheduling
- peer

Specify a peer partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): none
Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.

7.2.5.7 processors_per_iterator

- Keywords Area
- method
- multi_start
- processors_per_iterator

Specify the number of processors per iterator server when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional processors_per_iterator specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterations. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.

7.2.6 pareto_set

- Keywords Area
- method
- pareto_set

Pareto set optimization

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

In the pareto set minimization method (pareto_set), a series of optimization or least squares calibration runs are performed for different weightings applied to multiple objective functions. This set of optimal solutions defines a "Pareto set," which is useful for investigating design trade-offs between competing objectives. The code is similar enough to the multi_start technique that both algorithms are implemented in the same ConcurrentMetaIterator class.

The pareto_set specification must identify an optimization or least squares calibration method using either a method_pointer or a method_name plus optional model_pointer. This minimizer is responsible for computing a set of optimal solutions from a set of response weightings (multi-objective weights or least squares term weights). These weightings can be specified as follows: (1) using random_weight_sets, in which case weightings are selected randomly within [0,1] bounds, (2) using weight_sets, in which the weighting sets are specified in a list, or (3) using both random_weight_sets and weight_sets, for which the combined set of weights will be used. In aggregate, at least one set of weights must be specified. The set of optimal solutions is called the "pareto set," which can provide valuable design trade-off information when there are competing objectives.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the best parameters and responses returned by each sub-iterator. The weights are provided as metadata. See the Multistart and Pareto Set documentation for details.
7.2 method

- **method_name**

Specify sub-method by name

**Specification**

**Alias:** opt_method_name  
**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

The `method_name` keyword is used to specify a sub-method by Dakota method name (e.g. ‘npsol_sqp’) rather than block pointer. The method will be executed using its default settings. The optional `model_pointer` specification can be used to associate a model block with the method.

7.2.6.1.1 `model_pointer`

- **Keywords Area**
  - method
  - pareto_set
  - method_name
  - model_pointer

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

**Alias:** opt_model_pointer  
**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The `model_pointer` is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.
Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```dakota
environment
  tabular_data
  method_pointer = 'UQ'
method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
  sample_type lhs
  distribution cumulative
model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic
method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2
model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'
interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.6.2 method_pointer

  • Keywords Area
  • method
  • pareto_set
```
7.2 method

- **method_pointer**

Pointer to optimization or least-squares sub-method

**Topics**

This keyword is related to the topics:
- **block_pointer**

**Specification**

**Alias:** opt_method_pointer

**Argument(s):** STRING

**Description**

The `method_pointer` keyword is used to specify a pointer to an optimization or least-squares sub-method that is responsible for computing a set of optimal solutions for a set of response weightings.

### 7.2.6.3 random_weight_sets

- **Keywords Area**
- **method**
- **pareto_set**
- **random_weight_sets**

Number of random weighting sets

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
</tbody>
</table>

**Description**

The `pareto_set` specification must identify an optimization or least squares calibration method using either a `method_pointer` or a `method_name` plus optional `model_pointer`. This minimizer is responsible for computing a set of optimal solutions from a set of response weightings (multi-objective weights or least squares term weights). These weightings can be specified as follows: (1) using `random_weight_sets`, in which case weightings are selected randomly within [0,1] bounds, (2) using `weight_sets`, in which the weighting sets are specified in a list, or (3) using both `random_weight_sets` and `weight_sets`, for which the combined set of weights will be used. In aggregate, at least one set of weights must be specified. The set of optimal solutions is called the "pareto set," which can provide valuable design trade-off information when there are competing objectives.
7.2.6.3.1 seed

- Keywords Area
- method
- pareto_set
- random_weight_sets
- seed

Seed of the random number generator

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.6.4 weight_sets

- Keywords Area
- method
- pareto_set
- weight_sets

List of user-specified weighting sets
7.2 method

Specification

Alias: multi_objective_weight_sets

Argument(s): REALLIST

Description

The pareto_set specification must identify an optimization or least squares calibration method using either a method_pointer or a method_name plus optional model_pointer. This minimizer is responsible for computing a set of optimal solutions from a set of response weightings (multi-objective weights or least squares term weights). These weightings can be specified as follows: (1) using random_weight_sets, in which case weightings are selected randomly within [0,1] bounds, (2) using weight_sets, in which the weighting sets are specified in a list, or (3) using both random_weight_sets and weight_sets, for which the combined set of weights will be used. In aggregate, at least one set of weights must be specified. The set of optimal solutions is called the "pareto set," which can provide valuable design trade-off information when there are competing objectives.

7.2.6.5 iterator_servers

• Keywords Area
  • method
  • pareto_set
  • iterator_servers

Specify the number of iterator servers when Dakota is run in parallel

Topics

This keyword is related to the topics:

• concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional iterator_servers specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.6.6 iterator_scheduling

• Keywords Area
  • method
Specify the scheduling of concurrent iterators when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>master</td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel iterator scheduling</td>
</tr>
</tbody>
</table>

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional `iterator_scheduling` specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, `hybrid`, `multi_start`, and `pareto_set` component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.2.6.6.1 master

- Keywords Area
- method
- pareto_set
- iterator_scheduling
- master

Specify a dedicated master partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism
7.2 method

Specification

Alias: none
Argument(s): none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the “master”) dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.

7.2.6.6.2 peer

• Keywords Area
• method
• pareto_set
• iterator_scheduling
• peer

Specify a peer partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

• concurrency_and_parallelism

Specification

Alias: none
Argument(s): none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.

7.2.6.7 processors_per_iterator

• Keywords Area
• method
• pareto_set
• processors_per_iterator

Specify the number of processors per iterator server when Dakota is run in parallel
Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional processors_per_iterator specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterations. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.

7.2.7 branch_and_bound

- Keywords Area
- method
- branch_and_bound

(Experimental Capability) Solves a mixed integer nonlinear optimization problem

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required</th>
<th>Description of Group Local Optimizer Selection (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>method_pointer</td>
<td>Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

The branch-and-bound optimization methods solves mixed integer nonlinear optimization problems. It does so by partitioning the parameter space according to some criteria along the integer or discrete variables. It then relaxes (i.e., treats all variables as continuous) the sub-problems created by the partitions and solves each sub-problem with a continuous nonlinear optimization method. Results of the sub-problems are combined in such a way that yields the solution to the original optimization problem.

Default Behavior

Branch-and-bound expects all discrete variables to be relaxable. If your problem has categorical or otherwise non-relaxable discrete variables, then this is not the optimization method you are looking for.

Expected Output

The optimal solution and associated parameters will be printed to the screen output.

Usage Tips

The user must choose a nonlinear optimization method to solve the sub-problems. We recommend choosing a method that would be chosen to solve a continuous problem that has similar form to the mixed integer problem.

Examples

```dakota
environment     
   method_pointer = 'BandB'

method          
   id_method = 'BandB'
   branch_and_bound
      output verbose
      method_pointer = 'SubNLP'

method          
   id_method = 'SubNLP'
   coliny_ea
      seed = 12345
      max_iterations = 100
      max_function_evaluations = 100

variables,      
   continuous_design = 3
   initial_point -1.0  1.5  2.0
   upper_bounds  10.0 10.0 10.0
   lower_bounds -10.0 -10.0 -10.0
   descriptors   'x1' 'x2' 'x3'
   discrete_design_range = 2
   initial_point 2 2
   lower_bounds  1 1
   upper_bounds  4 9
   descriptors   'y1' 'y2'

interface,      
   fork
      analysis_driver = 'text_book'

responses,      
   objective_functions = 1
```
7.2.7.1 method_pointer

- Keywords Area
- method
- branch_and_bound
- method_pointer

Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Description

The method_pointer keyword is used to specify a pointer to an optimization or least-squares sub-method to apply in the context of what could be described as hierarchical methods. In surrogate-based methods, the sub-method is applied to the surrogate model. In the branch-and-bound method, the sub-method is applied to the relaxed sub-problems.

Any model_pointer identified in the sub-method specification is ignored. Instead, the parent method is responsible for selecting the appropriate model to use as specified by its model_pointer. In surrogate-based methods, it is a surrogate model defined using its model_pointer. In branch-and-bound methods, it is the relaxed model that is constructed internally from the original model.

7.2.7.2 method_name

- Keywords Area
- method
- branch_and_bound
- method_name

Specify sub-method by name

Specification

Alias: none

Argument(s): STRING

Child Keywords:
### Description

The `method_name` keyword is used to specify a sub-method by Dakota method name (e.g. `npsol_sqp`) rather than block pointer. The method will be executed using its default settings. The optional `model_pointer` specification can be used to associate a model block with the method.

#### 7.2.7.2.1 model_pointer

- **Keywords Area**
- **method**
- **branch_and_bound**
- **method_name**
- **model_pointer**

Identifier for model block to be used by a method

#### Topics

This keyword is related to the topics:

- **block_pointer**

#### Specification

- **Alias**: none
- **Argument(s)**: STRING
- **Default**: method use of last model parsed (or use of default model if none parsed)

#### Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

#### Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

#### Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.7.3 scaling

- Keywords Area
- method
- branch_and_bound
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `scales` keyword, which gives characteristic values (divisors)
- a `scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. log - logarithmic scaling
   First, any characteristic values from the optional `_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.
Scales

The `scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto`, `log` - optional
- `value` - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```dakota
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.8 surrogate_based_local

- Keywords Area
- method
- surrogate_based_local

Local Surrogate Based Optimization

Topics

This keyword is related to the topics:

- surrogate_based_optimization_methods

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Subproblem Optimizer Selection (Group 1)</td>
<td>method_pointer</td>
<td>Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>method_name</td>
<td>Specify sub-method by name</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>soft_convergence_limit</td>
<td>Limit number of iterations with little improvement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>truth_surrogate_bypass</td>
<td>Bypass lower level surrogates when performing truth verifications on a top level surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>approx_subproblem</td>
<td>Identify functions to be included in surrogate merit function</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>merit_function</td>
<td>Select type of penalty or merit function</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>acceptance_logic</td>
<td>Set criteria for trusted surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>constraint_relax</td>
<td>Enable constraint relaxation</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>trust_region</td>
<td>Specification group for trust region model management</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
</tbody>
</table>
Description

In surrogate-based optimization (SBO) and surrogate-based nonlinear least squares (SBNLS), minimization occurs using a set of one or more approximations, defined from a surrogate model, that are built and periodically updated using data from a “truth” model. The surrogate model can be a global data fit (e.g., regression or interpolation of data generated from a design of computer experiments), a multipoint approximation, a local Taylor Series expansion, or a model hierarchy approximation (e.g., a low-fidelity simulation model), whereas the truth model involves a high-fidelity simulation model. The goals of surrogate-based methods are to reduce the total number of truth model simulations and, in the case of global data fit surrogates, to smooth noisy data with an easily navigated analytic function.

In the surrogate-based local method, a trust region approach is used to manage the minimization process to maintain acceptable accuracy between the surrogate model and the truth model (by limiting the range over which the surrogate model is trusted). The process involves a sequence of minimizations performed on the surrogate model and bounded by the trust region. At the end of each approximate minimization, the candidate optimum point is validated using the truth model. If sufficient decrease has been obtained in the truth model, the trust region is re-centered around the candidate optimum point and the trust region will either shrink, expand, or remain the same size depending on the accuracy with which the surrogate model predicted the truth model decrease. If sufficient decrease has not been attained, the trust region center is not updated and the entire trust region shrinks by a user-specified factor. The cycle then repeats with the construction of a new surrogate model, a minimization, and another test for sufficient decrease in the truth model. This cycle continues until convergence is attained.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

Theory

For surrogate_based_local problems with nonlinear constraints, a number of algorithm formulations exist as described in[23] and as summarized in the Advanced Examples section of the Models chapter of the Users Manual[5].

See Also

These keywords may also be of interest:

- efficient_global
- surrogate_based_global
7.2.8.1 method_pointer

- Keywords Area
- method
- surrogate_based_local
- method_pointer

Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: approx_method_pointer

Argument(s): STRING

Description

The method_pointer keyword is used to specify a pointer to an optimization or least-squares sub-method to apply in the context of what could be described as hierarchical methods. In surrogate-based methods, the sub-method is applied to the surrogate model. In the branch-and-bound method, the sub-method is applied to the relaxed sub-problems.

Any model_pointer identified in the sub-method specification is ignored. Instead, the parent method is responsible for selecting the appropriate model to use as specified by its model_pointer. In surrogate-based methods, it is a surrogate model defined using its model_pointer. In branch-and-bound methods, it is the relaxed model that is constructed internally from the original model.

7.2.8.2 method_name

- Keywords Area
- method
- surrogate_based_local
- method_name

Specify sub-method by name

Specification

 Alias: approx_method_name
 Argument(s): STRING
Description

The `method_name` keyword is used to specify a sub-method by Dakota method name (e.g., 'npsol_sqp') rather than block pointer. The method will be executed using its default settings. The optional `model_pointer` specification can be used to associate a model block with the method.

7.2.8.3 `model_pointer`

- Keywords Area
- `method`
- `surrogate_based_local`
- `model_pointer`

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- `block_pointer`

Specification

**Alias:** approx_model_pointer

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
  samples = 10
```
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                   0.1 0.2 0.6
                   0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.8.4 soft_convergence_limit

- **Keywords Area**
- **method**
- **surrogate_based_local**
- **soft_convergence_limit**

Limit number of iterations w/ little improvement

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 5

**Description**

soft_convergence_limit (a soft convergence control for the surrogate_based_local iterations which limits the number of consecutive iterations with improvement less than the convergence tolerance). soft_convergence > 1
counts an iteration in which a design point was rejected as an iteration with no improvement. Setting soft_convergence=1 ignores iterations involving rejected points, instead comparing only consecutive iterations of accepted points.

7.2.8.5 truth_surrogate_bypass

- Keywords Area
- method
- surrogate_based_local
- truth_surrogate_bypass

Bypass lower level surrogates when performing truth verifications on a top level surrogate

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no bypass

**Description**

truth_surrogate_bypass (a flag for bypassing all lower level surrogates when performing truth verifications on a top level surrogate).

7.2.8.6 approx_subproblem

- Keywords Area
- method
- surrogate_based_local
- approx_subproblem

Identify functions to be included in surrogate merit function

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** original_primary original_constraints

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Objective Formulation (Group 1)</td>
<td>original_primary</td>
<td>Construct approximations of all primary functions</td>
</tr>
</tbody>
</table>
### Description

First, the "primary" functions (that is, the objective functions or calibration terms) in the approximate subproblem can be selected to be surrogates of the original primary functions (original_primary), a single objective function (single_objective) formed from the primary function surrogates, or either an augmented Lagrangian merit function (augmented_lagrangian_objective) or a Lagrangian merit function (lagrangian_objective) formed from the primary and secondary function surrogates. The former option may imply the use of a nonlinear least squares method, a multiobjective optimization method, or a single objective optimization method to solve the approximate subproblem, depending on the definition of the primary functions. The latter three options all imply the use of a single objective optimization method regardless of primary function definition. Second, the surrogate constraints in the approximate subproblem can be selected to be surrogates of the original constraints (original_constraints) or linearized approximations to the surrogate constraints (linearized_constraints), or constraints can be omitted from the subproblem (no_constraints).

#### 7.2.8.6.1 original_primary

- **Keywords Area**
- **method**
- **surrogate_based_local**
- **approx_subproblem**
- **original_primary**

Construct approximations of all primary functions

### Specification

**Alias**: none

**Argument(s)**: none

### Description

For SBL problems with nonlinear constraints, a number of algorithm formulations exist as described in[23] and as summarized in the Advanced Examples section of the Models chapter of the Users Manual[5]. First, the "primary"
functions (that is, the objective functions or calibration terms) in the approximate subproblem can be selected to be surrogates of the original primary functions (original_primary), a single objective function (single_objective) formed from the primary function surrogates, or either an augmented Lagrangian merit function (augmented_lagrangian_objective) or a Lagrangian merit function (lagrangian_objective) formed from the primary and secondary function surrogates. The former option may imply the use of a nonlinear least squares method, a multiobjective optimization method, or a single objective optimization method to solve the approximate subproblem, depending on the definition of the primary functions. The latter three options all imply the use of a single objective optimization method regardless of primary function definition.

7.2.8.6.2 single_objective

- Keywords Area
- method
- surrogate_based_local
- approx_subproblem
- single_objective

Construct approximation a single objective functions only

Specification

Alias: none
Argument(s): none

Description

For SBL problems with nonlinear constraints, a number of algorithm formulations exist as described in[23] and as summarized in the Advanced Examples section of the Models chapter of the Users Manual[5]. First, the "primary" functions (that is, the objective functions or calibration terms) in the approximate subproblem can be selected to be surrogates of the original primary functions (original_primary), a single objective function (single_objective) formed from the primary function surrogates, or either an augmented Lagrangian merit function (augmented_lagrangian_objective) or a Lagrangian merit function (lagrangian_objective) formed from the primary and secondary function surrogates. The former option may imply the use of a nonlinear least squares method, a multiobjective optimization method, or a single objective optimization method to solve the approximate subproblem, depending on the definition of the primary functions. The latter three options all imply the use of a single objective optimization method regardless of primary function definition.

7.2.8.6.3 augmented_lagrangian_objective

- Keywords Area
- method
- surrogate_based_local
- approx_subproblem
- augmented_lagrangian_objective

Augmented Lagrangian approximate subproblem formulation

Specification

Alias: none
Argument(s): none
Description

For SBL problems with nonlinear constraints, a number of algorithm formulations exist as described in[23] and as summarized in the Advanced Examples section of the Models chapter of the Users Manual[5]. First, the “primary” functions (that is, the objective functions or calibration terms) in the approximate subproblem can be selected to be surrogates of the original primary functions (original_primary), a single objective function (single_objective) formed from the primary function surrogates, or either an augmented Lagrangian merit function (augmented_lagrangian_objective) or a Lagrangian merit function (lagrangian_objective) formed from the primary and secondary function surrogates. The former option may imply the use of a nonlinear least squares method, a multiobjective optimization method, or a single objective optimization method to solve the approximate subproblem, depending on the definition of the primary functions. The latter three options all imply the use of a single objective optimization method regardless of primary function definition.

7.2.8.6.4 lagrangian_objective

- Keywords Area
- method
- surrogate_based_local
- approx_subproblem
- lagrangian_objective

Lagrangian approximate subproblem formulation

Specification

Alias: none
Argument(s): none

Description

For SBL problems with nonlinear constraints, a number of algorithm formulations exist as described in[23] and as summarized in the Advanced Examples section of the Models chapter of the Users Manual[5]. First, the “primary” functions (that is, the objective functions or calibration terms) in the approximate subproblem can be selected to be surrogates of the original primary functions (original_primary), a single objective function (single_objective) formed from the primary function surrogates, or either an augmented Lagrangian merit function (augmented_lagrangian_objective) or a Lagrangian merit function (lagrangian_objective) formed from the primary and secondary function surrogates. The former option may imply the use of a nonlinear least squares method, a multiobjective optimization method, or a single objective optimization method to solve the approximate subproblem, depending on the definition of the primary functions. The latter three options all imply the use of a single objective optimization method regardless of primary function definition.

7.2.8.6.5 original_constraints

- Keywords Area
- method
- surrogate_based_local
- approx_subproblem
- original_constraints

Use the constraints directly
### linearized_constraints

- **Keywords Area**
- **method**
- **surrogate_based_local**
- **approx_subproblem**
- **linearized_constraints**

Use linearized approximations to the constraints

### no_constraints

- **Keywords Area**
- **method**
- **surrogate_based_local**
- **approx_subproblem**
- **no_constraints**

Don’t use constraints
Description

The surrogate constraints in the approximate subproblem can be selected to be surrogates of the original constraints (original_constraints) or linearized approximations to the surrogate constraints (linearized_constraints), or constraints can be omitted from the subproblem (no_constraints).

7.2.8.7 merit_function

- Keywords Area
- method
- surrogate_based_local
- merit_function

Select type of penalty or merit function

Specification

Alias: none

Argument(s): none

Default: augmented_lagrangian_merit

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>penalty_merit</td>
<td>Use penalty merit function</td>
<td></td>
<td></td>
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<tr>
<td>adaptive_penalty_merit</td>
<td>Use adaptive penalty merit function</td>
<td></td>
<td></td>
</tr>
<tr>
<td>lagrangian_merit</td>
<td>Use first-order Lagrangian merit function</td>
<td></td>
<td></td>
</tr>
<tr>
<td>augmented_lagrangian_merit</td>
<td>Use combined penalty and zeroth-order Lagrangian merit function</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Following optimization of the approximate subproblem, the candidate iterate is evaluated using a merit function, which can be selected to be a simple penalty function with penalty ramped by surrogate_based_local iteration number (penalty_merit), an adaptive penalty function where the penalty ramping may be accelerated in order to avoid rejecting good iterates which decrease the constraint violation (adaptive_penalty_merit), a Lagrangian merit function which employs first-order Lagrange multiplier updates (lagrangian_merit), or an augmented Lagrangian merit function which employs both a penalty parameter and zeroth-order Lagrange multiplier updates (augmented_lagrangian_merit). When an augmented Lagrangian is selected for either the subproblem objective or the merit function (or both), updating of penalties and multipliers follows the approach described in[15].

7.2.8.7.1 penalty_merit

- Keywords Area
Second, the surrogate constraints in the approximate subproblem can be selected to be surrogates of the original constraints (original_constraints) or linearized approximations to the surrogate constraints (linearized_constraints), or constraints can be omitted from the subproblem (no_constraints). Following optimization of the approximate subproblem, the candidate iterate is evaluated using a merit function, which can be selected to be a simple penalty function with penalty ramped by SBL iteration number (penalty_merit), an adaptive penalty function where the penalty ramping may be accelerated in order to avoid rejecting good iterates which decrease the constraint violation (adaptive_penalty_merit), a Lagrangian merit function which employs first-order Lagrange multiplier updates (lagrangian_merit), or an augmented Lagrangian merit function which employs both a penalty parameter and zeroth-order Lagrange multiplier updates (augmented_lagrangian_merit). When an augmented Lagrangian is selected for either the subproblem objective or the merit function (or both), updating of penalties and multipliers follows the approach described in[15].
an adaptive penalty function where the penalty ramping may be accelerated in order to avoid rejecting good iterates which decrease the constraint violation (adaptive_penalty_merit), a Lagrangian merit function which employs first-order Lagrange multiplier updates (lagrangian_merit), or an augmented Lagrangian merit function which employs both a penalty parameter and zeroth-order Lagrange multiplier updates (augmented_lagrangian_merit). When an augmented Lagrangian is selected for either the subproblem objective or the merit function (or both), updating of penalties and multipliers follows the approach described in[15].

7.2.8.7.3 lagrangian_merit

- Keywords Area
- method
- surrogate_based_local
- merit_function
- lagrangian_merit

Use first-order Lagrangian merit function

Specification

Alias: none
Argument(s): none

Description

Second, the surrogate constraints in the approximate subproblem can be selected to be surrogates of the original constraints (original_constraints) or linearized approximations to the surrogate constraints (linearized_constraints), or constraints can be omitted from the subproblem (no_constraints). Following optimization of the approximate subproblem, the candidate iterate is evaluated using a merit function, which can be selected to be a simple penalty function with penalty ramped by SBL iteration number (penalty_merit), an adaptive penalty function where the penalty ramping may be accelerated in order to avoid rejecting good iterates which decrease the constraint violation (adaptive_penalty_merit), a Lagrangian merit function which employs first-order Lagrange multiplier updates (lagrangian_merit), or an augmented Lagrangian merit function which employs both a penalty parameter and zeroth-order Lagrange multiplier updates (augmented_lagrangian_merit). When an augmented Lagrangian is selected for either the subproblem objective or the merit function (or both), updating of penalties and multipliers follows the approach described in[15].

7.2.8.7.4 augmented_lagrangian_merit

- Keywords Area
- method
- surrogate_based_local
- merit_function
- augmented_lagrangian_merit

Use combined penalty and zeroth-order Lagrangian merit function

Specification

Alias: none
Argument(s): none
Second, the surrogate constraints in the approximate subproblem can be selected to be surrogates of the original constraints (original_constraints) or linearized approximations to the surrogate constraints (linearized_constraints), or constraints can be omitted from the subproblem (no_constraints). Following optimization of the approximate subproblem, the candidate iterate is evaluated using a merit function, which can be selected to be a simple penalty function with penalty ramped by SBL iteration number (penalty_merit), an adaptive penalty function where the penalty ramping may be accelerated in order to avoid rejecting good iterates which decrease the constraint violation (adaptive_penalty_merit), a Lagrangian merit function which employs first-order Lagrange multiplier updates (lagrangian_merit), or an augmented Lagrangian merit function which employs both a penalty parameter and zeroth-order Lagrange multiplier updates (augmented_lagrangian_merit). When an augmented Lagrangian is selected for either the subproblem objective or the merit function (or both), updating of penalties and multipliers follows the approach described in [15].

7.2.8.8 acceptance_logic

- Keywords Area
- method
- surrogate_based_local
- acceptance_logic

Set criteria for trusted surrogate

Specification

Alias: none

Argument(s): none

Default: filter

Child Keywords:

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<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Acceptance Logic (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>tr_ratio</td>
<td>Surrogate-Based Local iterate acceptance logic</td>
</tr>
</tbody>
</table>

Description

Following calculation of the merit function for the new iterate, the iterate is accepted or rejected and the trust region size is adjusted for the next surrogate_based_local iteration. Iterate acceptance is governed either by a trust region ratio (tr_ratio) formed from the merit function values or by a filter method (filter); however, trust region resizing logic is currently based only on the trust region ratio. For infeasible iterates, constraint relaxation can be used for balancing constraint satisfaction and progress made toward an optimum.

7.2.8.8.1 tr_ratio

- Keywords Area
- method
- surrogate_based_local
Surrogate-Based Local iterate acceptance logic

Specification

Alias: none
Argument(s): none

Description

Following calculation of the merit function for the new iterate, the iterate is accepted or rejected and the trust region size is adjusted for the next SBL iteration. Iterate acceptance is governed either by a trust region ratio \((tr\_ratio)\) formed from the merit function values or by a filter method \((filter)\); however, trust region resizing logic is currently based only on the trust region ratio. For infeasible iterates, constraint relaxation can be used for balancing constraint satisfaction and progress made toward an optimum. The command \texttt{constraint\_relax} followed by a method name specifies the type of relaxation to be used. Currently, homotopy\cite{71} is the only available method for constraint relaxation, and this method is dependent on the presence of the NPSOL library within the Dakota executable.

7.2.8.8.2 filter

• Keywords Area
• method
• surrogate\_based\_local
• acceptance\_logic
• filter

Surrogate-Based Local iterate acceptance logic

Specification

Alias: none
Argument(s): none

Description

Following calculation of the merit function for the new iterate, the iterate is accepted or rejected and the trust region size is adjusted for the next SBL iteration. Iterate acceptance is governed either by a trust region ratio \((tr\_ratio)\) formed from the merit function values or by a filter method \((filter)\); however, trust region resizing logic is currently based only on the trust region ratio. For infeasible iterates, constraint relaxation can be used for balancing constraint satisfaction and progress made toward an optimum. The command \texttt{constraint\_relax} followed by a method name specifies the type of relaxation to be used. Currently, homotopy\cite{71} is the only available method for constraint relaxation, and this method is dependent on the presence of the NPSOL library within the Dakota executable.
7.2.8.9  constraint_relax

- Keywords Area
- method
- surrogate_based_local
- constraint_relax

Enable constraint relaxation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no relaxation

**Child Keywords:**

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<tr>
<th>Required/Optional Required</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>homotopy</td>
<td>Surrogate-Based local constraint relaxation method for infeasible iterates</td>
</tr>
</tbody>
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**Description**

The command `constraint_relax` followed by a method name specifies the type of relaxation to be used. Currently, `homotopy` [71] is the only available method for constraint relaxation, and this method is dependent on the presence of the NPSOL library within the Dakota executable.

7.2.8.9.1  homotopy

- Keywords Area
- method
- surrogate_based_local
- constraint_relax
- homotopy

Surrogate-Based local constraint relaxation method for infeasible iterates

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Currently, `homotopy` [71] is the only available method for constraint relaxation, and this method is dependent on the presence of the NPSOL library within the Dakota executable.
7.2.8.10 trust_region

- Keywords Area
- method
- surrogate_based_local
- trust_region

Specification group for trust region model management

Specification

Alias: none
Argument(s): none

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Optional</td>
<td>initial_size</td>
<td>initial_size</td>
<td>Trust region initial size (relative to bounds)</td>
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<tr>
<td>Optional</td>
<td>minimum_size</td>
<td>minimum_size</td>
<td>Trust region minimum size</td>
</tr>
<tr>
<td>Optional</td>
<td>contract_threshold</td>
<td>contract_threshold</td>
<td>Shrink trust region if trust region ratio is below this value</td>
</tr>
<tr>
<td>Optional</td>
<td>expand_threshold</td>
<td>expand_threshold</td>
<td>Expand trust region if trust region ratio is above this value</td>
</tr>
<tr>
<td>Optional</td>
<td>contraction_factor</td>
<td>contraction_factor</td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td>expansion_factor</td>
<td>expansion_factor</td>
<td>Trust region expansion factor</td>
</tr>
</tbody>
</table>

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next surrogate_based_local iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next surrogate_based_local iteration.

Each of these specifications are REAL scalars, with the exception of initial_size, which is a REALLIST.

7.2.8.10.1 initial_size

- Keywords Area
Trust region initial size (relative to bounds)

**Specification**

**Alias:** none  
**Argument(s):** REALIST

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

All of these specifications are REAL scalars, with the exception of the specification of `initial_size`, which is a REALIST. This array corresponds to the case when there are more than 2 model forms or discretizations within a model hierarchy. The default `initial_size` involves a recursive halving of the global bounds for each trust region in the hierarchy: e.g., a scalar value of .5 for a single trust region managing two model forms/discretizations, or an array of (.125, .25, .5 ) for four model forms/discretizations (three trust regions ordered from the lowest to highest fidelity surrogate, with model four as truth).

7.2.8.10.2  `minimum_size`

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.e-6

Trust region minimum size
7.2 method

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.8.10.3 contract_threshold

- Keywords Area
- method
- surrogate_based_local
- trust_region
- contract_threshold

Shrink trust region if trust region ratio is below this value

Specification

Alias: none

Argument(s): REAL

Default: 0.25

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.8.10.4 expand_threshold

- Keywords Area
- method
- surrogate_based_local
- trust_region
- expand_threshold
Expand trust region if trust region ratio is above this value

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.75

**Description**

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.8.10.5 contraction_factor

- Keywords Area
- method
- surrogate_based_local
- trust_region
- contraction_factor

Amount by which step length is rescaled

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.25

**Description**

For pattern search methods, contraction_factor specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is $1 / \text{contraction\_factor}$

7.2.8.10.6 expansion_factor

- Keywords Area
- method
- surrogate_based_local
7.2 method

- trust_region
- expansion_factor

Trust region expansion factor

Specification

Alias: none
Argument(s): REAL
Default: 2.0

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.8.11 max_iterations

- Keywords Area
- method
- surrogate_based_local
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.8.12 convergence\_tolerance

- Keywords Area
- method
- surrogate\_based\_local
- convergence\_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method\_independent\_controls

Specification

\textbf{Alias}: none  
\textbf{Argument(s)}: REAL  
\textbf{Default}: 1.e-4

Description

The \texttt{convergence\_tolerance} specification provides a real value for controlling the termination of iteration.

For optimization, it is most commonly a \textbf{relative convergence tolerance} for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by \texttt{convergence\_tolerance}, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.8.13 constraint_tolerance

- Keywords Area
- method
- surrogate_based_local
- constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local: 1.0e-4
7.2.9 surrogate_based_global

- Keywords Area
- method
- surrogate_based_global

Global Surrogate Based Optimization

Topics

This keyword is related to the topics:

- surrogate_based_optimization_methods

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Required (Choose One)</td>
<td>Sub-method Selection (Group 1)</td>
<td>method_pointer</td>
<td>Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>method_name</td>
<td>Specify sub-method by name</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>replace_points</td>
<td>(Recommended) Replace points in the surrogate training set, instead of appending</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
</tbody>
</table>

Description

The `surrogate_based_global` specification must identify:

- a sub-method, using either `method_pointer` or `method_name`
7.2 method

- `model_pointer` must be used to identify a surrogate model

**surrogate_based_global** works in an iterative scheme where optimization is performed on a global surrogate using the same bounds during each iteration.

- In one iteration, the optimal solutions of the surrogate model are found, and then a selected set of these optimal surrogate solutions are passed to the next iteration.
- At the next iteration, these surrogate points are evaluated with the "truth" model, and then these points are added back to the set of points upon which the next surrogate is constructed.

In this way, the optimization acts on a more accurate surrogate during each iteration, presumably driving to optimality quickly.

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

**Method Independent Controls**

- `max_iterations` is used as a stopping criterion (see note below)

**Notes**

We have some cautionary notes before using the surrogate-based global method:

- **This approach has no guarantee of convergence.**
  - One might first try a single minimization method coupled with a surrogate model prior to using the surrogate-based global method. This is essentially equivalent to setting `max_iterations` to 1 and will allow one to get a sense of what surrogate types are the most accurate to use for the problem.
  - Also note that one can specify that surrogates be built for all primary functions and constraints or for only a subset of these functions and constraints. This allows one to use a "truth" model directly for some of the response functions, perhaps due to them being much less expensive than other functions.
  - We initially recommend a small number of maximum iterations, such as 3-5, to get a sense of how the optimization is evolving as the surrogate gets updated. If it appears to be changing significantly, then a larger number (used in combination with restart) may be needed.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- **Best Parameters**
- **Best Objective Functions** (when `objective_functions` are specified)
- **Best Nonlinear Constraints**
- **Calibration** (when `calibration_terms` are specified)

**Theory**

In surrogate-based optimization (SBO) and surrogate-based nonlinear least squares (SBNLS), minimization occurs using a set of one or more approximations, defined from a surrogate model, that are built and periodically updated using data from a "truth" model. The surrogate model can be a global data fit (e.g., regression or interpolation of data generated from a design of computer experiments), a multipoint approximation, a local Taylor Series expansion, or a
model hierarchy approximation (e.g., a low-fidelity simulation model), whereas the truth model involves a high-fidelity simulation model. The goals of surrogate-based methods are to reduce the total number of truth model simulations and, in the case of global data fit surrogates, to smooth noisy data with an easily navigated analytic function.

It was originally designed for MOGA (a multi-objective genetic algorithm). Since genetic algorithms often need thousands or tens of thousands of points to produce optimal or near-optimal solutions, the use of surrogates can be helpful for reducing the truth model evaluations. Instead of creating one set of surrogates for the individual objectives and running the optimization algorithm on the surrogate once, the idea is to select points along the (surrogate) Pareto frontier, which can be used to supplement the existing points.

In this way, one does not need to use many points initially to get a very accurate surrogate. The surrogate becomes more accurate as the iterations progress.

See Also

These keywords may also be of interest:

- efficient_global
- surrogate_based_local

7.2.9.1 method_pointer

- Keywords Area
- method
- surrogate_based_global
- method_pointer

Pointer to sub-method to apply to a surrogate or branch-and-bound sub-problem

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: approx_method_pointer

Argument(s): STRING

Description

The method_pointer keyword is used to specify a pointer to an optimization or least-squares sub-method to apply in the context of what could be described as hierarchical methods. In surrogate-based methods, the sub-method is applied to the surrogate model. In the branch-and-bound method, the sub-method is applied to the relaxed sub-problems.

Any model_pointer identified in the sub-method specification is ignored. Instead, the parent method is responsible for selecting the appropriate model to use as specified by its model_pointer. In surrogate-based methods, it is a surrogate model defined using its model_pointer. In branch-and-bound methods, it is the relaxed model that is constructed internally from the original model.
7.2 method

7.2.9.2 method_name

- Keywords Area
- method
- surrogate_based_global
- method_name

Specify sub-method by name

Specification

Alias: approx_method_name

Argument(s): STRING

Description

The method_name keyword is used to specify a sub-method by Dakota method name (e.g. 'npsol_sqp') rather than block pointer. The method will be executed using its default settings. The optional model_pointer specification can be used to associate a model block with the method.

7.2.9.3 model_pointer

- Keywords Area
- method
- surrogate_based_global
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: approx_model_pointer

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior
If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

### Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

### Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
       0.1 0.2 0.6
       0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

### 7.2.9.4 replace_points

- Keywords Area
- method
- surrogate_based_global
- replace_points
(Recommended) Replace points in the surrogate training set, instead of appending

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** Points appended, not replaced

**Description**

The user has the option of appending the optimal points from the surrogate model to the current set of truth points or using the optimal points from the surrogate model to replace the optimal set of points from the previous iteration. Although appending to the set is the default behavior, at this time we strongly recommend using the option `replace_points` because it appears to be more accurate and robust.

**7.2.9.5 max_iterations**

- **Keywords Area**  
- **method**  
- **surrogate_based_global**  
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed `max_iterations` iterations. In cases where there are multiple nested iterations to be controlled, `max_iterations` generally controls the outer-most context.

**Default Behavior**

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.10 dot_frcg

- Keywords Area
- method
- dot_frcg

DOT conjugate gradient optimization method

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td></td>
<td>Compute speculative gradients</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
</tbody>
</table>
### Description

DOT library\[88\] implementation of Fletcher-Reeves conjugate gradient method for unconstrained optimization. See package_dot for information common to all DOT methods.

**Caution regarding dot_frcg.** In DOT Version 4.20, we have noticed inconsistent behavior of this algorithm across different versions of Linux. Our best assessment is that it is due to different treatments of uninitialized variables. As we do not know the intention of the code authors and maintaining DOT source code is outside of the Dakota project scope, we have not made nor are we recommending any code changes to address this. However, all users who use dot_frcg in DOT Version 4.20 should be aware that results may not be reliable.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Calibration (when calibration_terms are specified)

### 7.2.10.1 max_iterations

- Keywords Area
  - method
  - dot_frcg
  - max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

### Topics

This keyword is related to the topics:

- method_independent_controls

### Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior
Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.10.2 convergence_tolerance

- Keywords Area
- method
- dot_frcg
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.10.3 constraint_tolerance

Keywords Area

method

dot_frcg

constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4
7.2.10.4 speculative

- Keywords Area
- method
- dot_frgc
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).
7.2.10.5 max_function_evaluations

• Keywords Area
• method
• dot_frcg
• max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.10.6 scaling

• Keywords Area
• method
• dot_frcg
• scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

• method_independent_controls
Specification

**Alias:** none

**Argument(s):** none

**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling

   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling

   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

   - two-sided bounds scaled into the interval $[0,1]$;
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. **log** - logarithmic scaling

   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.
7.2 method

Scales
The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples
The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.10.7 model_pointer

- Keywords Area
- method
- dot_frcg
- model_pointer

Identifier for model block to be used by a method

Topics
This keyword is related to the topics:

- block_pointer

Specification
Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

environment
    tabular_data
method_pointer = 'UQ'

method
    id_method = 'UQ'
model_pointer = 'SURR'
sampling,
    samples = 10
    seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
distribution cumulative

model
    id_model = 'SURR'
surrogate global,
    dace_method_pointer = 'DACE'
polynomial quadratic

method
    id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
    id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians
7.2.11 dot_mmfd

- Keywords Area
- method
- dot_mmfd

DOT modified method of feasible directions

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td>constraint_-tolerance</td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td>max_function_-evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

DOT library[88] implementation of the modified method of feasible directions for constrained optimization.

See package_dot for information common to all DOT methods.
DOT requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- **Best Parameters**
- **Best Objective Functions** (when objective functions are specified)
- **Best Nonlinear Constraints**
- **Calibration** (when calibration terms are specified)

### 7.2.11.1 max_iterations

- **Keywords Area**
- **method**
- **dot_mmfd**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- **max_function_evaluations**
7.2.11.2  convergence_tolerance

- Keywords Area
- method
- dot_mmfd
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.11.3  constraint_tolerance

- Keywords Area
- method
Maximum allowable constraint violation still considered feasible

Topics
This keyword is related to the topics:

- method_independent_controls

Specification

_Alias_: none
_Argument(s)_: REAL
_Default_: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

_Defaults_

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

7.2.11.4 speculative

- Keywords Area
- method
- dot_mmfd
- speculative

Compute speculative gradients

Topics
This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.11.5 max_function_evaluations

- Keywords Area
- method
- dot_mmfd
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description
The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed `max_function_evaluations` evaluations. See also `max_iterations`.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

Default Behavior
Default value is 1000.

7.2.11.6  scaling

- Keywords Area
- method
- dot_mmfd
- scaling

Turn on scaling for variables, responses, and constraints

Topics
This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description
Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.
Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the `variables` and `responses` sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval $[0,1]$;
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. **log** - logarithmic scaling
   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.
   - Logarithmic scaling is not available for linear constraints.
   - When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The `*scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `*scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto`, `log` - optional
- `value` - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.11.7 model_pointer

- Keywords Area
- method
- dot_mmf
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

- **Alias:** none
- **Argument(s):** STRING
- **Default:** method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.
7.2 method

Examples

```dakota
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
  samples = 10
  seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
  0.1 0.2 0.6
  0.1 0.2 0.6
  sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
  lower_bounds = 0.0.
  upper_bounds = 1.0.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
response_functions = 3
  no_gradients
  no_hessians
```

7.2.12 dot_bfgs

- Keywords Area
- method
- dot_bfgs

DOT BFGS optimization method

Specification

Alias: none
Argument(s): none
Child Keywords:
## Description

DOT library[88] implementation of Broyden-Fletcher-Goldfarb-Shanno method for unconstrained optimization. See package_dot for information common to all DOT methods. 

*DOT requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives.*

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- **Best Parameters**
- **Best Objective Functions** (when objective_functions) are specified)
- **Calibration** (when calibration_terms are specified)

### 7.2.12.1 max_iterations

- **Keywords Area**
7.2 method

- method
- dot_bfgs
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: $25+n$)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed $\text{max\_iterations}$ iterations. In cases where there are multiple nested iterations to be controlled, $\text{max\_iterations}$ generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max_function_evaluations

7.2.12.2 convergence_tolerance

- Keywords Area
- method
- dot_bfgs
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

**Notes on each library:**

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See `nl2sol`
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.12.3 `constraint_tolerance`

- **Keywords Area**
  - method
  - `dot_bfgs`
  - `constraint_tolerance`

Maximum allowable constraint violation still considered feasible

**Topics**

This keyword is related to the topics:

- `method_independent_controls`

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** Library default
Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint’s violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

7.2.12.4 speculative

- Keywords Area
  - method
  - dot_bfgs
  - speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative
gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.12.5 max_function_evaluations

- Keywords Area
- method
- dot_bfgs
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.
7.2.12.6 scaling

- Keywords Area
- method
- dot_bfgs
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.
2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval \([0,1]\);
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into \([0,1]\).

3. **log** - logarithmic scaling
   First, any characteristic values from the optional \(^*_scales\) specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The \(^*_scales\) keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding \(^*_scale_types\), as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0e10\*DBL_MIN\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.12.7 **model_pointer**

- **Keywords Area**
- **method**
- **dot_bfgs**
- **model_pointer**

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.13 dot_slp

- Keywords Area
- method
- dot_slp

DOT Sequential Linear Program

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
</tbody>
</table>
### Description

DOT library[88] implementation of sequential linear programming for constrained optimization.

See package_dot for information common to all DOT methods.

DOT requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

#### 7.2.13.1 max_iterations

- Keywords Area
- method
- dot_slp
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25∗n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max_iterations} \) generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- `max_function_evaluations`

**7.2.13.2 convergence_tolerance**

- **Keywords Area**
- **method**
- **dot_slp**
- **convergence_tolerance**

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The **convergence_tolerance** specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.13.3 constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.
Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

7.2.13.4 speculative

- Keywords Area
- method
- dot_slp
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search
in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.13.5 max_function_evaluations

- Keywords Area
- method
- dot_slp
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.13.6 scaling

- Keywords Area
- method
- dot_slp
- scaling

Turn on scaling for variables, responses, and constraints
This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `∗scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `∗scales` keyword, which gives characteristic values (divisors)  
- a `∗scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling  
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling  
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];  
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;  
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].
3. log-logarithmic scaling

First, any characteristic values from the optional *scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10\times\text{DBL_MIN}$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.13.7 model_pointer

- Keywords Area
- method
- dot_slp
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer
Specifications

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
    sample_type lhs
distribution cumulative
model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2
model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
7.2 method

    analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.14  dot_sqp

    • Keywords Area

    • method

    • dot_sqp

DOT Sequential Quadratic Program

Topics

This keyword is related to the topics:

    • package_dot

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>constraint_-tolerance</td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>speculative</td>
<td>Compute speculative gradients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_function_-evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

DOT library\[88\] implementation of sequential quadratic programming for constrained optimization.

See package_dot for information common to all DOT methods.

*DOT requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives.*

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

7.2.14.1 max_iterations

- Keywords Area
Number of iterations allowed for optimizers and adaptive UQ methods

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \( \text{max\_function\_evaluations} \)

7.2.14.2 convergence_tolerance

Keywords Area
- method
- dot_sqp
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- \( \text{method\_independent\_controls} \)
Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do. Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.14.3 `constraint_tolerance`

- Keywords Area
- method
- dot_sqp
- constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: Library default
Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint’s violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

7.2.14.4 speculative

- Keywords Area
- method
- dot_sqp
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative
gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to \texttt{responses} gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

### 7.2.14.5 max\_function\_evaluations

- **Keywords Area**
- **method**
- **dot\_sqp**
- **max\_function\_evaluations**

Number of function evaluations allowed for optimizers

**Topics**

This keyword is related to the topics:

- **method\_independent\_controls**

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max\_function\_evaluations} evaluations. See also \texttt{max\_iterations}.

Some optimizers (e.g., \texttt{ncsu\_direct}) may run past this limit in the course of an iteration step that began before \texttt{max\_function\_evaluations} was exceeded.

**Default Behavior**

Default value is 1000.
7.2.14.6 scaling

- Keywords Area
  - method
  - dot_sqp
  - scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.
2. auto - automatic scaling

   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   • two-sided bounds scaled into the interval [0,1];
   • one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   • no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. log - logarithmic scaling

   First, any characteristic values from the optional *scales specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

   • auto, log - optional
   • value - required

   A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```dakota
responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = "value"
   primary_scales = 1 1 100

responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = "value" "value" "value"
   primary_scales = 1 1 100
```

7.2.14.7 model_pointer

   • Keywords Area
   • method
   • dot_sqp
   • model_pointer

   Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
    method_pointer = ‘UQ’

method
  id_method = ‘UQ’
  model_pointer = ‘SURR’
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = ‘SURR’
    surrogate global,
    dace_method_pointer = ‘DACE’
    polynomial quadratic

method
  id_method = ‘DACE’
  model_pointer = ‘DACE_M’
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = ‘DACE_M’
  single
  interface_pointer = ‘I1’
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ’x1’ ’x2’

interface
id_interface = ’I1’
system asynch evaluation_concurrency = 5
analysis_driver = ’text_book’

responses
response_functions = 3
no_gradients
no_hessians

7.2.15 conmin_frcg

• Keywords Area
• method
• conmin_frcg

CONMIN conjugate gradient optimization method

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
</tbody>
</table>
Optional \hspace{1cm} speculative

Optional \hspace{1cm} max_function_evaluations

Optional \hspace{1cm} scaling

Optional \hspace{1cm} model_pointer

### Description

The interpretations of the method independent controls for CONMIN are essentially identical to those for DOT. See package_dot for information related to CONMIN methods, specifically dot_frcg.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified)
- Calibration (when calibration_terms are specified)

### See Also

These keywords may also be of interest:

- dot_frcg

#### 7.2.15.1 max_iterations

- Keywords Area
- method
- conmin_frcg
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

### Topics

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+\(n\))

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \(\text{max_iterations}\) iterations. In cases where there are multiple nested iterations to be controlled, \(\text{max_iterations}\) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \text{max_function_evaluations}

7.2.15.2 convergence_tolerance

- Keywords Area
- method
- conmin_frcg
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- \text{methodIndependent_controls}

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The \text{convergence_tolerance} specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.15.3 constraint tolerance

- Keywords Area
- method
- conmin_frcg
- constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.
Defaults

Defaults vary depending on the method.

- CONMIN optimizers: $1.0\times 10^{-3}$
- DOT constrained optimizers: $3.0\times 10^{-3}$
- surrogate_based_local : $1.0\times 10^{-4}$

7.2.15.4 speculative

- Keywords Area
- method
- conmin_frcg
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search
in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.15.5 max_function_evaluations

- Keywords Area
- method
- conmin_frcg
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.15.6 scaling

- Keywords Area
- method
- conmin_frcg
- scaling

Turn on scaling for variables, responses, and constraints
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].
3. log-logarithmic scaling

First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```plaintext
responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value"
    primary_scales = 1 1 100

responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value" "value" "value"
    primary_scales = 1 1 100
```

7.2.15.7 model_pointer

- Keywords Area
- method
- conmin_frcg
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer
Specification

**Alias:** none  

**Argument(s):** STRING  

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```plaintext
evironment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121
  seed = 5034

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system async evaluation_concurrency = 5
```
7.2 method

    analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.16 conmin_mfd

- Keywords Area
- method
- conmin_mfd

CONMIN method of feasible directions

Topics

This keyword is related to the topics:

- package_conmin

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>---------------</td>
<td>-------------------------------</td>
<td></td>
</tr>
<tr>
<td></td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The interpretations of the method independent controls for CONMIN are essentially identical to those for DOT.

See package_dot for information related to CONMIN methods, specifically dot_mmfd.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

**See Also**

These keywords may also be of interest:

- dot_mmfd

**7.2.16.1 max_iterations**

- Keywords Area
- method
- conmin_mfd
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls
7.2 method

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max_function_evaluations

---

7.2.16.2 convergence_tolerance

**Keywords Area**

- method
- conmin_mfd
- convergence_tolerance

**Description**

The convergence_tolerance specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a **relative convergence tolerance** for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- **DOT**: relative tolerance that must be satisfied for two consecutive iterations
- **NL2SOL**: See nl2sol
- **NLPQLP**: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- **NPSOL**: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.16.3 constraint_tolerance

**Keywords Area**

- **method**
- **conmin_mfd**
- **constraint_tolerance**

Maximum allowable constraint violation still considered feasible

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

<table>
<thead>
<tr>
<th>Alias:</th>
<th>none</th>
</tr>
</thead>
<tbody>
<tr>
<td>Argument(s):</td>
<td>REAL</td>
</tr>
<tr>
<td>Default:</td>
<td>Library default</td>
</tr>
</tbody>
</table>

**Description**

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.
 Defaults
Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

 7.2.16.4 speculative

- Keywords Area
- method
- conmin_mfd
- speculative

Compute speculative gradients

Topics
This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description
When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search
in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.16.5 max_function_evaluations

- Keywords Area
- method
- conmin_mfd
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.16.6 scaling

- Keywords Area
- method
- conmin_mfd
- scaling

Turn on scaling for variables, responses, and constraints
Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

• a *scales keyword, which gives characteristic values (divisors)
• a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   • two-sided bounds scaled into the interval [0,1];
   • one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   • no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Futher, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].
3. **log** - logarithmic scaling

   First, any characteristic values from the optional starscales specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
objective_functions 3
sense "maximize"
primary_scale_types = "value"
primary_scales = 1 1 100

responses
objective_functions 3
sense "maximize"
primary_scale_types = "value" "value" "value"
primary_scales = 1 1 100
```

**7.2.16.7 model_pointer**

- **Keywords Area**
- **method**
- **conmin_mfd**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**
7.2 method

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```dakota
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
      sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
```
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

### 7.2.17 dl_solver

- **Keywords Area**
- **method**
- **dl_solver**

(Experimental) Dynamically-loaded solver

**Topics**

This keyword is related to the topics:

- optimization_and_calibration

**Specification**

**Alias:** none

**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

This keyword specifies a dynamically-loaded optimization solver library, an experimental Dakota feature that is not enabled by default.

### 7.2.17.1 max_function_evaluations

- **Keywords Area**
- **method**
7.2 method

- dl_solver
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:
- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.17.2 scaling

- Keywords Area
- method
- dl_solver
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:
- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling
Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or ∗scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a ∗scales keyword, which gives characteristic values (divisors)
- a ∗scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. log - logarithmic scaling
   First, any characteristic values from the optional ∗_scales specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The ∗scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding ∗scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0 \times 10^{10}$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

### 7.2.17.3 model_pointer

- Keywords Area
- method
- dl_solver
- model_pointer

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- block_pointer

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

**Usage Tips**
When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians
```

7.2.18  `npsol_sqp`

- Keywords Area
- method
- npsol_sqp

NPSOL Sequential Quadratic Program
Topics

This keyword is related to the topics:

- package_npsol
- sequential_quadratic_programming
- local_optimization_methods

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>verify_level</td>
<td>verify_level</td>
<td>Verify the quality of analytic gradients</td>
</tr>
<tr>
<td>Optional</td>
<td>function_precision</td>
<td>function_precision</td>
<td>Specify the maximum precision of the analysis code responses</td>
</tr>
<tr>
<td>Optional</td>
<td>linesearch_-tolerance</td>
<td>linesearch_-tolerance</td>
<td>Choose how accurately the algorithm will compute the minimum in a line search</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td>constraint_-tolerance</td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
</tr>
</tbody>
</table>
### Description

Sequential quadratic programming optimizer.

*NPSOL requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives.*

### Stopping Criteria

The method independent controls for `max_iterations` and `max_function_evaluations` limit the number of major SQP iterations and the number of function evaluations that can be performed during an NPSOL optimization. The `convergence_tolerance` control defines NPSOL's internal optimality tolerance which is used in evaluating if an iterate satisfies the first-order Kuhn-Tucker conditions for a minimum. The magnitude of `convergence_tolerance` approximately specifies the number of significant digits of accuracy desired in the final objective function (e.g., `convergence_tolerance = 1.e-6` will result in approximately six digits of accuracy in the final objective function). The `constraint_tolerance` control defines how tightly the constraint functions are satisfied at convergence. The default value is dependent upon the machine precision of the platform in use, but is typically on the order of `1.e-8` for double precision computations. Extremely small values for `constraint_tolerance` may not be attainable. The `output verbosity` setting controls the amount of information generated at each major SQP iteration: the `silent` and `quiet` settings result in only one line of diagnostic output for each major iteration and print the final optimization solution, whereas the `verbose` and `debug` settings add additional information on the objective function, constraints, and variables at each major iteration.

### Concurrency

NPSOL is not a parallel algorithm and cannot directly take advantage of concurrent evaluations. However, if `numerical_gradients` with `method_source dakota` is specified, then the finite difference function evaluations can be performed concurrently (using any of the parallel modes described in the Users Manual [5]).

An important related observation is the fact that NPSOL uses two different line searches depending on how gradients are computed. For either `analytic_gradients` or `numerical_gradients with method_source dakota`, NPSOL is placed in user-supplied gradient mode (NPSOL's "Derivative Level" is set to 3) and it uses a gradient-based line search (the assumption is that user-supplied gradients are inexpensive). On the other hand, if `numerical_gradients` are selected with `method_source vendor`, then NPSOL is computing finite differences internally and it will use a value-based line search (the assumption is that finite differencing on each line search evaluation is too expensive). The ramifications of this are: (1) performance will vary between `method_source dakota` and `method_source vendor`, and (2) gradient speculation is unnecessary when performing optimization in parallel since the gradient-based line search in user-supplied gradient mode is already load balanced for parallel execution. Therefore, a speculative specification will be ignored by NPSOL, and optimization with numerical gradients should select `method_source dakota` for load balanced parallel operation and `method_source vendor` for efficient serial operation.

### Linear constraints

Lastly, NPSOL supports specialized handling of linear inequality and equality constraints. By specifying the coefficients and bounds of the linear inequality constraints and the coefficients and targets of the linear equality constraints, this information can be provided to NPSOL at initialization and tracked internally, removing the need for the user to provide the values of the linear constraints on every function evaluation.
Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

7.2.18.1 verify_level

- Keywords Area
- method
- npsol_sqp
- verify_level

Verify the quality of analytic gradients

Specification

Alias: none
Argument(s): INTEGER
Default: -1 (no gradient verification)

Description

verify_level instructs the NPSOL and NLSSOL algorithms to perform their own finite difference verification of the gradients provided by Dakota. Typically these are used to verify analytic_gradients produced by a simulation code, though the option can be used with other Dakota-supplied gradient types including numerical or mixed.

Level 1 will verify the objective gradients, level 2, the nonlinear constraint gradients, and level 3, both. See the Optional Input Parameters section of the NPSOL manual[33] for additional information, including options to verify at the user-supplied initial point vs. first feasible point.

7.2.18.2 function_precision

- Keywords Area
- method
- npsol_sqp
- function_precision

Specify the maximum precision of the analysis code responses
Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0e-10

**Description**

The `function_precision` control provides the algorithm with an estimate of the accuracy to which the problem functions can be computed. This is used to prevent the algorithm from trying to distinguish between function values that differ by less than the inherent error in the calculation.

### 7.2.18.3 `linesearch_tolerance`

- **Keywords Area**
- **method**
- **npsol_sqp**
- **linesearch_tolerance**

Choose how accurately the algorithm will compute the minimum in a line search

### Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0.9 (inaccurate line search)

**Description**

The `linesearch_tolerance` setting controls the accuracy of the line search. The smaller the value (between 0 and 1), the more accurately the algorithm will attempt to compute a precise minimum along the search direction.

### 7.2.18.4 `convergence_tolerance`

- **Keywords Area**
- **method**
- **npsol_sqp**
- **convergence_tolerance**

Stopping criterion based on objective function convergence

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4
7.2 method

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

NPSOL defines an internal optimality tolerance which is used in evaluating if an iterate satisfies the first-order Kuhn-Tucker conditions for a minimum. The magnitude of convergence_tolerance approximately specifies the number of significant digits of accuracy desired in the final objective function (e.g., convergence_tolerance = 1.0e-6 will result in approximately six digits of accuracy in the final objective function).

7.2.18.5 max_iterations

• Keywords Area
  • method
  • npsol_sqp
  • max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

  • method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

  • max_function_evaluations
7.2.18.6  constraint_tolerance

- Keywords Area
- method
- npsol_sqp
- constraint_tolerance

Maximum allowable constraint violation still considered feasible

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** Library default

**Description**

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint’s violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Dakota’s `constraint_tolerance` specification maps to NPSOL’s feasibility tolerance which is an absolute tolerance on the constraints.

7.2.18.7  speculative

- Keywords Area
- method
- npsol_sqp
- speculative

Compute speculative gradients

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no speculation
When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

### 7.2.18.8 max_function_evaluations

- **Keywords Area**
- **method**
- **npsol_sqp**
- **max_function_evaluations**

Number of function evaluations allowed for optimizers

### Topics

This keyword is related to the topics:

- **method_independent_controls**

### Specification

- **Alias:** none
- **Argument(s):** INTEGER
- **Default:** 1000
Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max\_function\_evaluations} evaluations. See also \texttt{max\_iterations}.

Some optimizers (e.g. \texttt{ncsu\_direct}) may run past this limit in the course of an iteration step that began before \texttt{max\_function\_evaluations} was exceeded.

Default Behavior

Default value is 1000.

7.2.18.9 scaling

- Keywords Area
- method
- npsol_sqp
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- \texttt{method\_independent\_controls}

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the \texttt{scaling} keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords \texttt{scale\_types} and/or \texttt{\ast scales} in the \texttt{variables} and \texttt{responses} blocks. When the \texttt{scaling} keyword is omitted from \texttt{method}, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a \texttt{\ast scales} keyword, which gives characteristic values (divisors)
- a \texttt{\ast scale\_types} keyword, which determines how to use the characteristic values
The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints. When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```
7.2.18.10  model_pointer

- Keywords Area
- method
- npsol_sqp
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
  dace_method_pointer = 'DACE'
polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.19 nlssol_sqp

  • Keywords Area
  • method
  • nlssol_sqp

Sequential Quadratic Program for nonlinear least squares

Topics

This keyword is related to the topics:

  • sequential_quadratic_programming
  • nonlinear_least_squares

Specification

Alias: none

Argument(s): none

Child Keywords:
### Optional

- **function_precision**
  - Specify the maximum precision of the analysis code responses.

- **linesearch_tolerance**
  - Choose how accurately the algorithm will compute the minimum in a line search.

- **convergence_tolerance**
  - Stopping criterion based on objective function convergence.

- **max_iterations**
  - Number of iterations allowed for optimizers and adaptive UQ methods.

- **constraint_tolerance**
  - Maximum allowable constraint violation still considered feasible.

- **speculative**
  - Compute speculative gradients.

- **max_function_evaluations**
  - Number of function evaluations allowed for optimizers.

- **scaling**
  - Turn on scaling for variables, responses, and constraints.

- **model_pointer**
  - Identifier for model block to be used by a method.

### Description

NLSSOL supports unconstrained, bound-constrained, and generally-constrained least-squares calibration problems. It exploits the structure of a least squares objective function through the periodic use of Gauss-Newton Hessian approximations to accelerate the SQP algorithm.

**NLSSOL requires a separate software license and therefore may not be available in all versions of Dakota.** nl2sol or optpp_g_newton may be suitable alternatives.

### Stopping Criteria

The method independent controls for `max_iterations` and `max_function_evaluations` limit the number of major SQP iterations and the number of function evaluations that can be performed during an NPSOL optimization. The `convergence_tolerance` control defines NPSOL's internal optimality tolerance which is used in evaluating if an iterate satisfies the first-order Kuhn-Tucker conditions for a minimum. The magnitude of `convergence_tolerance` approximately specifies the number of significant digits of accuracy desired in the optimization.
final objective function (e.g., convergence_tolerance $= 1.e-6$ will result in approximately six digits of accuracy in the final objective function). The constraint_tolerance control defines how tightly the constraint functions are satisfied at convergence. The default value is dependent upon the machine precision of the platform in use, but is typically on the order of $1.e-8$ for double precision computations. Extremely small values for constraint_tolerance may not be attainable.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)
- Parameter Confidence Intervals (when num_experiments equals 1)

**See Also**

These keywords may also be of interest:

- npsol_sqp
- nl2sol
- optpp_g_newton
- field_calibration_terms

### 7.2.19.1 verify_level

- **Keywords Area**
- method
- nlssol_sqp
- verify_level

Verify the quality of analytic gradients

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** -1 (no gradient verification)

**Description**

`verify_level` instructs the NPSOL and NLSSOL algorithms to perform their own finite difference verification of the gradients provided by Dakota. Typically these are used to verify analytic_gradients produced by a simulation code, though the option can be used with other Dakota-supplied gradient types including numerical or mixed.

Level 1 will verify the objective gradients, level 2, the nonlinear constraint gradients, and level 3, both. See the Optional Input Parameters section of the NPSOL manual[33] for additional information, including options to verify at the user-supplied initial point vs. first feasible point.
7.2.19.2  function_precision

Specify the maximum precision of the analysis code responses

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0e-10

**Description**

The `function_precision` control provides the algorithm with an estimate of the accuracy to which the problem functions can be computed. This is used to prevent the algorithm from trying to distinguish between function values that differ by less than the inherent error in the calculation.

7.2.19.3  linesearch_tolerance

Choose how accurately the algorithm will compute the minimum in a line search

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.9 (inaccurate line search)

**Description**

The `linesearch_tolerance` setting controls the accuracy of the line search. The smaller the value (between 0 and 1), the more accurately the algorithm will attempt to compute a precise minimum along the search direction.

7.2.19.4  convergence_tolerance
7.2 method

- nlssol_sqp
- convergence_tolerance

Stopping criterion based on objective function convergence

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

NPSOL defines an internal optimality tolerance which is used in evaluating if an iterate satisfies the first-order Kuhn-Tucker conditions for a minimum. The magnitude of convergence_tolerance approximately specifies the number of significant digits of accuracy desired in the final objective function (e.g., convergence_tolerance = 1.0e-6 will result in approximately six digits of accuracy in the final objective function).

7.2.19.5 max_iterations

- Keywords Area
- method
- nlssol_sqp
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \( \text{max\_function\_evaluations} \)

7.2.19.6 \hspace{1em} \text{constraint\_tolerance}

Maximum allowable constraint violation still considered feasible

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint’s violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Dakota’s constraint\_tolerance settings maps to NPSOL’s feasibility tolerance which is an absolute tolerance on the constraints.

7.2.19.7 \hspace{1em} \text{speculative}

Compute speculative gradients
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.19.8 max_function_evaluations

- Keywords Area
- method
- nlssol_sqp
- max_function_evaluations

Number of function evaluations allowed for optimizers
Topics
This keyword is related to the topics:

- `method_independent_controls`

Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed `max_function_evaluations` evaluations. See also `max_iterations`.

Some optimizers (e.g. `ncsu_direct`) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

**Default Behavior**

Default value is 1000.

7.2.19.9 scaling

- `Keywords Area`  
- `method`  
- `nlssol_sqp`  
- `scaling`

Turn on scaling for variables, responses, and constraints

Topics
This keyword is related to the topics:

- `method_independent_controls`

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- A `*scales` keyword, which gives characteristic values (divisors)
- A `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

### Scale Types

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The `*scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `*scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto, log` - optional
- `value` - required

A warning is issued if scaling would result in division by a value smaller in magnitude than `1.0e10*DBL_MIN`. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value"
    primary_scales = 1 1 100

responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value" "value" "value"
    primary_scales = 1 1 100
```

7.2.19.10 model_pointer

- Keywords Area
- method
- nlssol_sqp
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.
7.2 method

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
  samples = 10
  seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
  sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.20 nlpql_sqp

• Keywords Area
• method
  • nlpql_sqp

NLPQL Sequential Quadratic Program

Topics

This keyword is related to the topics:

• package_nlpql
**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td></td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

NLPQL implementation of sequential quadratic programming. The particular SQP implementation in `nlpql_sqp` uses a variant with distributed and non-monotone line search. Thus, this variant is designed to be more robust in the presence of inaccurate or noisy gradients common in many engineering applications.

_NLPQL requires a separate software license and therefore may not be available in all versions of Dakota. CONMIN or OPT++ methods may be suitable alternatives._

The method independent controls for maximum iterations and output verbosity are mapped to NLPQL controls MAX-IT and IPRINT, respectively. The maximum number of function evaluations is enforced within the NLPQLPOptimizer class.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
• Best Nonlinear Constraints
• Calibration (when calibration_terms are specified)

7.2.20.1 max_iterations

• Keywords Area
• method
• nlpql_sqp
• max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations

7.2.20.2 convergence_tolerance

• Keywords Area
• method
• nlpql_sqp
• convergence_tolerance

Stopping criterion based on objective function or statistics convergence
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

 Argument(s): REAL

 Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.20.3 max_function_evaluations

- Keywords Area
- method
- nlpql_sqp
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls
7.2 method

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed `max_function_evaluations` evaluations. See also `max_iterations`.

Some optimizers (e.g. `ncsu_direct`) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

**Default Behavior**

Default value is 1000.

### 7.2.20.4 scaling

- **Keywords Area**
- **method**
- **nlpql_sqp**
- **scaling**

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.
Scaling also requires the specification of additional keywords `scale_types` and/or `*_scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*_scales` keyword, which gives characteristic values (divisors)
- a `*_scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component
   
   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The `*_scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `*_scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto, log` - optional
- `value` - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than `1.0e10*DBL_MIN`. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.20.5 model_pointer

- Keywords Area
- method
- nlpql_sqp
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.21 optpp_cg

• Keywords Area
• method
• optpp_cg

A conjugate gradient optimization method

Topics

This keyword is related to the topics:

• package_optpp
• local_optimization_methods
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_step</td>
<td>max_step</td>
<td>Max change in design point</td>
</tr>
<tr>
<td>Optional</td>
<td>gradient_tolerance</td>
<td>gradient_tolerance</td>
<td>Stopping criteria based on L2 norm of gradient</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td>max_function_-evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

The conjugate gradient method is an implementation of the Polak-Ribiere approach and handles only unconstrained problems.

See package_optpp for info related to all optpp methods.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- **Best Parameters**
- **Best Objective Functions** (when objective_functions are specified)
- **Calibration** (when calibration_terms are specified)
See Also

These keywords may also be of interest:

- optpp_g_newton
- optpp_pds
- optpp_fd_newton
- optpp_newton
- optpp_g_newton

7.2.21.1 max_step

- Keywords Area
- method
- optpp_cg
- max_step

Max change in design point

Specification

Alias: none
Argument(s): REAL
Default: 1000.

Description

The max_step control specifies the maximum step that can be taken when computing a change in the current design point (e.g., limiting the Newton step computed from current gradient and Hessian information). It is equivalent to a move limit or a maximum trust region size. The gradient_tolerance control defines the threshold value on the L2 norm of the objective function gradient that indicates convergence to an unconstrained minimum (no active constraints). The gradient_tolerance control is defined for all gradient-based optimizers.

7.2.21.2 gradient_tolerance

- Keywords Area
- method
- optpp_cg
- gradient_tolerance

Stopping criteria based on L2 norm of gradient

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4
Description

The \texttt{gradient\_tolerance} control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The \texttt{gradient\_tolerance} control is currently defined only for the OPT++ and ROL libraries.

7.2.21.3 \texttt{max\_iterations}

- Keywords Area
- method
- optpp\_cg
- \texttt{max\_iterations}

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- \texttt{method\_independent\_controls}

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: INTEGER

\textbf{Default}: 100 (exceptions: fsu\_cvt , local\_reliability: 25; global\_reliability , interval\_est , evidence) / efficient\_global: \(25 \times n\)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

\textbf{Default Behavior}

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.21.4 \texttt{convergence\_tolerance}

- Keywords Area
- method
• optpp_cg
• convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:
• method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

• DOT: relative tolerance that must be satisfied for two consecutive iterations
• NL2SOL: See nl2sol
• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

Compute speculative gradients
7.2 method

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.21.6 max_function_evaluations

- Keywords Area
- method
- optpp_cg
- max_function_evaluations

Number of function evaluations allowed for optimizers
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max\_function\_evaluations} evaluations. See also \texttt{max\_iterations}.

Some optimizers (e.g. \texttt{ncsu\_direct}) may run past this limit in the course of an iteration step that began before \texttt{max\_function\_evaluations} was exceeded.

Default Behavior

Default value is 1000.

7.2.21.7 scaling

- Keywords Area
- method
- optpp\_cg
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the \texttt{scaling} keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.21.8  model_pointer

- Keywords Area
- method
- optpp_cg
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.22 optpp_q_newton

- Keywords Area
- method
- optpp_q_newton

Quasi-Newton optimization method

Topics

This keyword is related to the topics:

- package_optpp
- local_optimization_methods
Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>search_method</td>
<td></td>
<td>Select a search method for Newton-based optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>merit_function</td>
<td></td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>steplength_to_boundary</td>
<td></td>
<td>Controls how close to the boundary of the feasible region the algorithm is allowed to move</td>
</tr>
<tr>
<td>Optional</td>
<td>centering_parameter</td>
<td></td>
<td>Controls how closely the algorithm should follow the &quot;central path&quot;</td>
</tr>
<tr>
<td>Optional</td>
<td>max_step</td>
<td></td>
<td>Max change in design point</td>
</tr>
<tr>
<td>Optional</td>
<td>gradient_tolerance</td>
<td></td>
<td>Stopping criteria based on L2 norm of gradient</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td></td>
<td>Compute speculative gradients</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
</tbody>
</table>
Description

This is a Newton method that expects a gradient and computes a low-rank approximation to the Hessian. Each of the Newton-based methods are automatically bound to the appropriate OPT++ algorithm based on the user constraint specification (unconstrained, bound-constrained, or generally-constrained). In the generally-constrained case, the Newton methods use a nonlinear interior-point approach to manage the constraints.

See package_optpp for info related to all optpp methods.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

7.2.22.1 search_method

- Keywords Area
- method
- optpp_q_newton
- search_method

Select a search method for Newton-based optimizers

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Line Search Method (Group 1)</td>
<td>value-based_line_search</td>
<td>Use only function values for line search</td>
</tr>
</tbody>
</table>
The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

### 7.2.22.1 `value_based_line_search`

- **Keywords Area**
  - `method`
  - `optpp_q_newton`
  - `search_method`
  - `value_based_line_search`

Use only function values for line search.

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** `trust_region` (unconstrained), `value_based_line_search` (bound/general constraints)

### Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems.
generally-constrained problems, and use of a \texttt{trust\_region} search method is the default for unconstrained problems.

\subsection{gradient\_based\_line\_search}

\begin{itemize}
\item Keywords Area
\item method
\item optpp\_q\_newton
\item search\_method
\item gradient\_based\_line\_search
\end{itemize}

Set the search method to use the gradient

\textbf{Specification}

\textbf{Alias:} none
\textbf{Argument(s):} none

\textbf{Description}

The \texttt{search\_method} control is defined for all Newton-based optimizers and is used to select between \texttt{trust\_region}, \texttt{gradient\_based\_line\_search}, and \texttt{value\_based\_line\_search} methods. The \texttt{gradient\_based\_line\_search} option uses the line search method proposed by\cite{63}. This option satisfies sufficient decrease and curvature conditions; whereas, \texttt{value\_based\_line\_search} only satisfies the sufficient decrease condition. At each line search iteration, the \texttt{gradient\_based\_line\_search} method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the \texttt{value\_based\_line\_search} method is preferred to the \texttt{gradient\_based\_line\_search} method. Each of these Newton methods additionally supports the \texttt{tr\_pds} selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a \texttt{trust\_region} search method is the default for unconstrained problems.

\subsection{trust\_region}

\begin{itemize}
\item Keywords Area
\item method
\item optpp\_q\_newton
\item search\_method
\item trust\_region
\end{itemize}

Use trust region as the globalization strategy.

\textbf{Specification}

\textbf{Alias:} none
\textbf{Argument(s):} none
Keywords Area

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.22.1.4 tr_pds

- Keywords Area
- method
- optpp_q_newton
- search_method
- tr_pds

Use direct search as the local search in a trust region method

Specification

Alias: none
Argument(s): none

Description

The search_method control is defined for all Newton-based optimizers and is used to select between trust_region, gradient_based_line_search, and value_based_line_search methods. The gradient_based_line_search option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, value_base_line_search only satisfies the sufficient decrease condition. At each line search iteration, the gradient_based_line_search method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the value_based_line_search method is preferred to the gradient_based_line_search method. Each of these Newton methods additionally supports the tr_pds selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a trust_region search method is the default for unconstrained problems.

7.2.22.2 merit_function

- Keywords Area
- method
- optpp_q_newton
- merit_function

Balance goals of reducing objective function and satisfying constraints
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** argaez_tapia

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
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<td><strong>Merit Function</strong> (Group 1)</td>
<td>el_bakry</td>
<td>El-Bakry merit function</td>
</tr>
<tr>
<td>argaez_tapia</td>
<td>The merit function by Tapia and Argaez</td>
<td></td>
<td></td>
</tr>
<tr>
<td>van_shanno</td>
<td>The merit function by Vanderbei and Shanno</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

A **merit_function** is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

7.2.22.2.1 el_bakry

- Keywords Area
- method
- optpp_q_newton
- merit_function
- el_bakry

El-Bakry merit function

Specification

**Alias:** none

**Argument(s):** none

Description

The "el_bakry" merit function is the L2-norm of the first order optimality conditions for the nonlinear programming problem. The cost per linesearch iteration is \( n+1 \) function evaluations. For more information, see[20].

7.2.22.2.2 argaez_tapia

- Keywords Area
- method
- optpp_q_newton
- merit_function
The merit function by Tapia and Argaez

Specification

Alias: none
Argument(s): none

Description

The "argaez_tapia" merit function can be classified as a modified augmented Lagrangian function. The augmented Lagrangian is modified by adding to its penalty term a potential reduction function to handle the perturbed complementarity condition. The cost per linesearch iteration is one function evaluation. For more information, see [83].

If the function evaluation is expensive or noisy, set the merit_function to "argaez_tapia" or "van_shanno".

7.2.22.2.3 van_shanno

• Keywords Area
• method
• optpp_q_newton
• merit_function
• van_shanno

The merit function by Vanderbei and Shanno

Specification

Alias: none
Argument(s): none

Description

The "van_shanno" merit function can be classified as a penalty function for the logarithmic barrier formulation of the nonlinear programming problem. The cost per linesearch iteration is one function evaluation. For more information see [85].

If the function evaluation is expensive or noisy, set the merit_function to "argaez_tapia" or "van_shanno".

7.2.22.3 steplength_to_boundary

• Keywords Area
• method
• optpp_q_newton
• steplength_to_boundary

Controls how close to the boundary of the feasible region the algorithm is allowed to move
7.2 method

Specification

Alias: none
Argument(s): REAL
Default: Merit function dependent: 0.8 (el_bakry), 0.99995 (argaez_tapia), 0.95 (van_shanno)

Description

The `steplength_to_boundary` specification is a parameter (between 0 and 1) that controls how close to the boundary of the feasible region the algorithm is allowed to move. A value of 1 means that the algorithm is allowed to take steps that may reach the boundary of the feasible region. If the user wishes to maintain strict feasibility of the design parameters this value should be less than 1. Default values are .8, .99995, and .95 for the `el_bakry`, `argaez_tapia`, and `van_shanno` merit functions, respectively.

7.2.22.4 centering_parameter

- Keywords Area
- method
- optpp_q_newton
- centering_parameter

Controls how closely the algorithm should follow the "central path"

Specification

Alias: none
Argument(s): REAL
Default: Merit function dependent: 0.2 (el_bakry), 0.2 (argaez_tapia), 0.1 (van_shanno)

Description

The `centering_parameter` specification is a parameter (between 0 and 1) that controls how closely the algorithm should follow the "central path". See[92] for the definition of central path. The larger the value, the more closely the algorithm follows the central path, which results in small steps. A value of 0 indicates that the algorithm will take a pure Newton step. Default values are .2, .2, and .1 for the `el_bakry`, `argaez_tapia`, and `van_shanno` merit functions, respectively.

7.2.22.5 max_step

- Keywords Area
- method
- optpp_q_newton
- max_step

Max change in design point
Specification

Alias: none
Argument(s): REAL
Default: 1000.

Description

The `max_step` control specifies the maximum step that can be taken when computing a change in the current design point (e.g., limiting the Newton step computed from current gradient and Hessian information). It is equivalent to a move limit or a maximum trust region size. The `gradient_tolerance` control defines the threshold value on the L2 norm of the objective function gradient that indicates convergence to an unconstrained minimum (no active constraints). The `gradient_tolerance` control is defined for all gradient-based optimizers.

7.2.22.6 gradient_tolerance

- Keywords Area
- method
- optpp_q_newton
- gradient_tolerance

Stopping criteria based on L2 norm of gradient

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The `gradient_tolerance` control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The `gradient_tolerance` control is currently defined only for the OPT++ and ROL libraries.

7.2.22.7 max_iterations

- Keywords Area
- method
- optpp_q_newton
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.22.8 convergence_tolerance

- Keywords Area
- method
- optpp_q_newton
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4
The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.22.9 speculative

- Keywords Area
- method
- optpp_q_newton
- speculative

Compute speculative gradients

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no speculation

**Description**

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based optimization...
optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a
trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint
violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted se-
quentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation
phase begins. By speculating that the gradient information associated with a given line search trial point will be
used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by
finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This
balances the total amount of computation to be performed at each design point and allows for efficient utilization
of multiple processors. While the total amount of work performed will generally increase (since some speculative
gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease
(since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel
during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for
the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical
or analytic gradient selections in the responses specification (refer to responses gradient section for information
on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite
difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also com-
puted speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search
in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced
parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where
applicable, for parallel optimization studies. By speculating that the derivative information at the current point will
be used later, the complete data set (all available gradient/Hessian information) can be computed on every function
evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load
balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism
in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.22.10 max_function_evaluations

- Keywords Area
- method
- optpp_q_newton
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached
any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations
evaluations. See also max_iterations.
Some optimizers (e.g., ncsu_direct) may run past this limit in the course of an iteration step that began before \texttt{max\_function\_evaluations} was exceeded.

**Default Behavior**

Default value is 1000.

7.2.22.11 scaling

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{optpp\_q\_newton}
- \texttt{scaling}

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- \texttt{method\_independent\_controls}

**Specification**

\textbf{Alias}: none

\textbf{Argument(s)}: none

\textbf{Default}: no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the \texttt{scaling} keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords \texttt{scale\_types} and/or \texttt{*scales} in the \texttt{variables} and \texttt{responses} blocks. When the \texttt{scaling} keyword is omitted from \texttt{method}, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a \texttt{*scales} keyword, which gives characteristic values (divisors)
- a \texttt{*scale\_types} keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in \texttt{responses} for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *scales specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

   - **auto, log** - optional
   - **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = *value*
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = *value* *value* *value*
  primary_scales = 1 1 100
```

**7.2.22.12 model_pointer**

- **Keywords Area**
- **method**
• optpp_a_newton
• model_pointer

Identifier for model block to be used by a method

Topics
This keyword is related to the topics:
• block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)

Description
The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior
If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips
When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.
See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
polynomial quadratic

method
  id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = ‘x1’ ‘x2’

interface
    id_interface = ’I1’
system asynch evaluation_concurrency = 5
    analysis_driver = ’text_book’

responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.23 optpp_fd_newton

    • Keywords Area
    • method
    • optpp_fd_newton

Finite Difference Newton optimization method

Topics

This keyword is related to the topics:

    • package_optpp
    • local_optimization_methods

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>search_method</td>
<td></td>
<td>Select a search method for Newton-based optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>merit_function</td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
<td></td>
</tr>
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<td>----------</td>
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<td>---------------------------------------------------------------------</td>
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<td>Optional</td>
<td>steplength_to_-boundary</td>
<td>Controls how close to the boundary of the feasible region the algorithm is allowed to move</td>
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<tr>
<td>Optional</td>
<td>centering_-parameter</td>
<td>Controls how closely the algorithm should follow the &quot;central path&quot;</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_step</td>
<td>Max change in design point</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gradient_tolerance</td>
<td>Stopping criteria based on L2 norm of gradient</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
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<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
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<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

This is a Newton method that expects a gradient and computes a finite-difference approximation to the Hessian. Each of the Newton-based methods are automatically bound to the appropriate OPT++ algorithm based on the user constraint specification (unconstrained, bound-constrained, or generally-constrained). In the generally-constrained case, the Newton methods use a nonlinear interior-point approach to manage the constraints.

See package_optpp for info related to all optpp methods.

**Expected HDF5 Output**
If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

See Also

These keywords may also be of interest:

- optpp_cg
- optpp_g_newton
- optpp_pds
- optpp_newton
- optpp_g_newton

7.2.23.1 search_method

- Keywords Area
- method
- optpp_fd_newton
- search_method

Select a search method for Newton-based optimizers

Specification

Alias: none

Argument(s): none

Child Keywords:

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<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Line Search Method (Group 1)</td>
<td>value_based_line_search</td>
<td>Use only function values for line search</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gradient_based_line_search</td>
<td>Set the search method to use the gradient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>trust_region</td>
<td>Use trust region as the globalization strategy</td>
</tr>
</tbody>
</table>
Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.23.1.1 `value_based_line_search`

- **Keywords Area**
- **method**
- **optpp_fd_newton**
- **search_method**
- **value_based_line_search**

Use only function values for line search

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** trust_region (unconstrained), value_based_line_search (bound/general constraints)

Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.23.1.2 `gradient_based_line_search`

- **Keywords Area**
7.2 method

- method
- optpp_fd_newton
- search_method
- gradient_based_line_search

Set the search method to use the gradient

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.23.1.3 trust_region

- Keywords Area
- method
- optpp_fd_newton
- search_method
- trust_region

Use trust region as the globalization strategy.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used...
when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.23.1.4 tr_pds

- **Keywords Area**
- **method**
- optpp_fd_newton
- search_method
- tr_pds

Use direct search as the local search in a trust region method

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.23.2 merit_function

- **Keywords Area**
- **method**
- optpp_fd_newton
- merit_function

Balance goals of reducing objective function and satisfying constraints
7.2 method

Specification

Alias: none

Argument(s): none

Default: argaez_tapia

Child Keywords:

<table>
<thead>
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<th>Dakota Keyword</th>
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<td>Merit Function (Group 1)</td>
<td>el_bakry</td>
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<tr>
<td>argaez_tapia</td>
<td>The merit function by Tapia and Argaez</td>
<td></td>
<td></td>
</tr>
<tr>
<td>van_shanno</td>
<td>The merit function by Vanderbei and Shanno</td>
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Description

A merit_function is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

7.2.23.2.1 el_bakry

- Keywords Area
- method
- optpp_fd_newton
- merit_function
- el_bakry

El-Bakry merit function

Specification

Alias: none

Argument(s): none

Description

The "el_bakry" merit function is the L2-norm of the first order optimality conditions for the nonlinear programming problem. The cost per linesearch iteration is $n+1$ function evaluations. For more information, see[20].

7.2.23.2.2 argaez_tapia

- Keywords Area
- method
- optpp_fd_newton
- merit_function
The merit function by Tapia and Argaez

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The "argaez_tapia" merit function can be classified as a modified augmented Lagrangian function. The augmented Lagrangian is modified by adding to its penalty term a potential reduction function to handle the perturbed complementarity condition. The cost per linesearch iteration is one function evaluation. For more information, see [83].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

7.2.23.2 van_shanno

- Keywords Area
- method
- optpp_fd_newton
- merit_function
- van_shanno

The merit function by Vanderbei and Shanno

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The "van_shanno" merit function can be classified as a penalty function for the logarithmic barrier formulation of the nonlinear programming problem. The cost per linesearch iteration is one function evaluation. For more information see [85].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

7.2.23.3 steplength_to_boundary

- Keywords Area
- method
- optpp_fd_newton
- steplength_to_boundary

Controls how close to the boundary of the feasible region the algorithm is allowed to move.
7.2 method

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** Merit function dependent: 0.8 (el_bakry), 0.99995 (argaez_tapia), 0.95 (van_shanno)

Description

The `steplength_to_boundary` specification is a parameter (between 0 and 1) that controls how close to the boundary of the feasible region the algorithm is allowed to move. A value of 1 means that the algorithm is allowed to take steps that may reach the boundary of the feasible region. If the user wishes to maintain strict feasibility of the design parameters this value should be less than 1. Default values are .8, .99995, and .95 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.23.4 `centering_parameter`

- **Keywords Area**
- **method**
- **optpp_fd_newton**
- **centering_parameter**

Controls how closely the algorithm should follow the "central path"

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** Merit function dependent: 0.2 (el_bakry), 0.2 (argaez_tapia), 0.1 (van_shanno)

Description

The `centering_parameter` specification is a parameter (between 0 and 1) that controls how closely the algorithm should follow the "central path". See[92] for the definition of central path. The larger the value, the more closely the algorithm follows the central path, which results in small steps. A value of 0 indicates that the algorithm will take a pure Newton step. Default values are .2, .2, and .1 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.23.5 `max_step`

- **Keywords Area**
- **method**
- **optpp_fd_newton**
- **max_step**

Max change in design point
Specification

Alias: none
Argument(s): REAL
Default: 1000.

Description

The max_step control specifies the maximum step that can be taken when computing a change in the current design point (e.g., limiting the Newton step computed from current gradient and Hessian information). It is equivalent to a move limit or a maximum trust region size. The gradient_tolerance control defines the threshold value on the L2 norm of the objective function gradient that indicates convergence to an unconstrained minimum (no active constraints). The gradient_tolerance control is defined for all gradient-based optimizers.

7.2.23.6 gradient_tolerance

- Keywords Area
- method
- optpp_fd_newton
- gradient_tolerance

Stopping criteria based on L2 norm of gradient

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The gradient_tolerance control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The gradient_tolerance control is currently defined only for the OPT++ and ROL libraries.

7.2.23.7 max_iterations

- Keywords Area
- method
- optpp_fd_newton
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods
7.2 method

Topics
This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description
The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior
Default value is 100.

See Also
These keywords may also be of interest:

- max_function_evaluations

7.2.23.8 convergence_tolerance

- Keywords Area
- method
- optpp_fd_newton
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics
This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4
The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.23.9 speculative

- Keywords Area
- method
- optpp_fd_newton
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based
optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a
trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint
violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted se-
quentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation
phase begins. By speculating that the gradient information associated with a given line search trial point will be
used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by
finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This
balances the total amount of computation to be performed at each design point and allows for efficient utilization
of multiple processors. While the total amount of work performed will generally increase (since some speculative
gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease
(since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel
during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for
the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical
or analytic gradient selections in the responses specification (refer to responses gradient section for information
on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite
difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also com-
puted speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search
in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced
parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where
applicable, for parallel optimization studies. By speculating that the derivative information at the current point will
be used later, the complete data set (all available gradient/Hessian information) can be computed on every function
evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load
balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism
in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.23.10 max_function_evaluations

- Keywords Area
  - method
  - optpp_fd_newton
  - max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached
any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations
evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

**Default Behavior**

Default value is 1000.

---

### 7.2.23.11 scaling

- **Keywords Area**
- **method**
- **optpp_fd_newton**
- **scaling**

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:
1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   • two-sided bounds scaled into the interval [0,1];
   • one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   • no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. log - logarithmic scaling
   First, any characteristic values from the optional \(*_scales\) specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

Scales
The \(*_scales\) keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding \(*_scale_types\), as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

   • auto, log - optional
   • value - required.

   A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples
The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.23.12 model_pointer

   • Keywords Area
   • method
• optpp_fd_newton
• model_pointer

Identifier for model block to be used by a method

Topics
This keyword is related to the topics:

• block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)

Description
The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior
If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips
When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.
See block_pointer for details about pointers.

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.24  optpp_g_newton

  • Keywords Area
  • method
  • optpp_g_newton

Newton method based least-squares calibration

Topics

This keyword is related to the topics:

  • package_optpp
  • local_optimization_methods

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>search_method</td>
<td>Select a search method for Newton-based optimizers</td>
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<tr>
<td>Optional</td>
<td>merit_function</td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>----------------</td>
<td>--------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>steplength_to_ - boundary</td>
<td>Controls how close to the boundary of the feasible region the algorithm is allowed to move</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>centering_ - parameter</td>
<td>Controls how closely the algorithm should follow the &quot;central path&quot;</td>
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<tr>
<td>Optional</td>
<td>max_step</td>
<td>Max change in design point</td>
<td></td>
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<td>gradient_tolerance</td>
<td>Stopping criteria based on L2 norm of gradient</td>
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<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
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<td>convergence_ - tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
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<td>Optional</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
<td></td>
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<td>Optional</td>
<td>max_function_ - evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
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<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
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</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
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</table>

**Description**

The Gauss-Newton algorithm is available as `optpp_g_newton` and supports unconstrained, bound-constrained, and generally-constrained problems. When interfaced with the unconstrained, bound-constrained, and nonlinear interior point full-Newton optimizers from the OPT++ library, it provides a Gauss-Newton least squares capability which – on zero-residual test problems – can exhibit quadratic convergence rates near the solution. (Real problems almost never have zero residuals, i.e., perfect fits.)

See `package_optpp` for info related to all `optpp` methods.
Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)
- Parameter Confidence Intervals (when num_experiments equals 1)

See Also

These keywords may also be of interest:

- optpp_cg
- optpp_pds
- optpp_fd_newton
- optpp_newton
- optpp_g_newton
- field_calibration_terms

7.2.24.1 search_method

Select a search method for Newton-based optimizers

Specification

Alias: none
Argument(s): none
Child Keywords: 

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<td>Line Search Method (Group 1)</td>
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<td>Use only function values for line search</td>
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<td></td>
<td></td>
<td>gradient_based_line_search</td>
<td>Set the search method to use the gradient</td>
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Dakota Version 6.15 Reference Manual generated on November 5, 2021
Use trust region as the globalization strategy.

Use direct search as the local search in a trust region method.

### Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_base_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

#### 7.2.24.1.1 value_based_line_search

- **Keywords Area**
  - `method`
  - `optpp_g_newton`
  - `search_method`
  - `value_based_line_search`

Use only function values for line search.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** `trust_region` (unconstrained), `value_based_line_search` (bound/general constraints)

### Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_base_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.
7.2.24.1.2 gradient-based-line-search

- Keywords Area
- method
- optpp_g_newton
- search_method
- gradient_based_line_search

Set the search method to use the gradient

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_base_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.24.1.3 trust_region

- Keywords Area
- method
- optpp_g_newton
- search_method
- trust_region

Use trust region as the globalization strategy.

**Specification**

**Alias:** none

**Argument(s):** none
Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.24.1.4 tr_pds

- Keywords Area
- method
- optpp_g_newton
- search_method
- tr_pds

Use direct search as the local search in a trust region method

Specification

Alias: none
Argument(s): none

Description

The search_method control is defined for all Newton-based optimizers and is used to select between trust_region, gradient_based_line_search, and value_based_line_search methods. The gradient_based_line_search option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, value_based_line_search only satisfies the sufficient decrease condition. At each line search iteration, the gradient_based_line_search method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the value_based_line_search method is preferred to the gradient_based_line_search method. Each of these Newton methods additionally supports the tr_pds selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a trust_region search method is the default for unconstrained problems.

7.2.24.2 merit_function

- Keywords Area
- method
- optpp_g_newton
- merit_function

Balance goals of reducing objective function and satisfying constraints
7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Default**: argaez_tapia

**Child Keywords**:

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<td><strong>el_bakry</strong></td>
<td>El-Bakry merit function</td>
</tr>
</tbody>
</table>

**Description**

A **merit function** is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

7.2.24.2.1 **el_bakry**

- **Keywords Area**
- **method**
- **optpp_g_newton**
- **merit_function**
- **el_bakry**

El-Bakry merit function

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The "el_bakry" merit function is the L2-norm of the first order optimality conditions for the nonlinear programming problem. The cost per linesearch iteration is \( n+1 \) function evaluations. For more information, see[20].

7.2.24.2.2 **argaez_tapia**

- **Keywords Area**
- **method**
- **optpp_g_newton**
- **merit_function**
The merit function by Tapia and Argaez

**Specification**

**Alias:** none  
**Argument(s):** none  

**Description**

The "argaez_tapia" merit function can be classified as a modified augmented Lagrangian function. The augmented Lagrangian is modified by adding to its penalty term a potential reduction function to handle the perturbed complementarity condition. The cost per linesearch iteration is one function evaluation. For more information, see [83].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

### 7.2.24.2.3 van_shanno

- Keywords Area  
- method  
- optpp_g_newton  
- merit_function  
- van_shanno

The merit function by Vanderbei and Shanno

**Specification**

**Alias:** none  
**Argument(s):** none  

**Description**

The "van_shanno" merit function can be classified as a penalty function for the logarithmic barrier formulation of the nonlinear programming problem. The cost per linesearch iteration is one function evaluation. For more information see[85].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

### 7.2.24.3 steplength_to_boundary

- Keywords Area  
- method  
- optpp_g_newton  
- steplength_to_boundary

Controls how close to the boundary of the feasible region the algorithm is allowed to move
7.2 method

Specification

Alias: none

Argument(s): REAL

Default: Merit function dependent: 0.8 (el_bakry), 0.99995 (argaez_tapia), 0.95 (van_shanno)

Description

The steplength_to_boundary specification is a parameter (between 0 and 1) that controls how close to the boundary of the feasible region the algorithm is allowed to move. A value of 1 means that the algorithm is allowed to take steps that may reach the boundary of the feasible region. If the user wishes to maintain strict feasibility of the design parameters this value should be less than 1. Default values are .8, .99995, and .95 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.24.4 centering_parameter

- Keywords Area
- method
- optpp_g_newton
- centering_parameter

Controls how closely the algorithm should follow the “central path”

Specification

Alias: none

Argument(s): REAL

Default: Merit function dependent: 0.2 (el_bakry), 0.2 (argaez_tapia), 0.1 (van_shanno)

Description

The centering_parameter specification is a parameter (between 0 and 1) that controls how closely the algorithm should follow the “central path”. See[92] for the definition of central path. The larger the value, the more closely the algorithm follows the central path, which results in small steps. A value of 0 indicates that the algorithm will take a pure Newton step. Default values are .2, .2, and .1 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.24.5 max_step

- Keywords Area
- method
- optpp_g_newton
- max_step

Max change in design point
Specification

Alias: none
Argument(s): REAL
Default: 1000.

Description

The \texttt{max\_step} control specifies the maximum step that can be taken when computing a change in the current design point (e.g., limiting the Newton step computed from current gradient and Hessian information). It is equivalent to a move limit or a maximum trust region size. The \texttt{gradient\_tolerance} control defines the threshold value on the L2 norm of the objective function gradient that indicates convergence to an unconstrained minimum (no active constraints). The \texttt{gradient\_tolerance} control is defined for all gradient-based optimizers.

7.2.24.6 \texttt{gradient\_tolerance}

\begin{itemize}
  \item Keywords Area
  \item method
  \item optpp\_g\_newton
  \item gradient\_tolerance
\end{itemize}

Stopping critiera based on L2 norm of gradient

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The \texttt{gradient\_tolerance} control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The \texttt{gradient\_tolerance} control is currently defined only for the OPT++ and ROL libraries.

7.2.24.7 \texttt{max\_iterations}

\begin{itemize}
  \item Keywords Area
  \item method
  \item optpp\_g\_newton
  \item max\_iterations
\end{itemize}

Number of iterations allowed for optimizers and adaptive UQ methods
7.2 method

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25∗n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \textit{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \textit{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max\_function\_evaluations

7.2.24.8 convergence_tolerance

- Keywords Area
- method
- optpp\_g\_newton
- convergence\_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4
Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.24.9 speculative

- Keywords Area
- method
- optpp_g_newton
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based
optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.24.10 max_function_evaluations

- Keywords Area
- method
- optpp_g_newton
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

**Default Behavior**

Default value is 1000.

### 7.2.24.11 scaling

- Keywords Area
- method
- optpp_g_newton
- scaling

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `scales` in the variables and responses blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `scales` keyword, which gives characteristic values (divisors)
- a `scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:
1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   • two-sided bounds scaled into the interval [0,1];
   • one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   • no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. log - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

   • auto, log - optional
   • value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```plaintext
responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = *value*
   primary_scales = 1 1 100

responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = *value* *value* *value*
   primary_scales = 1 1 100
```

7.2.24.12 model_pointer

   • Keywords Area
   • method
Keywords

- `optpp_g_newton`
- `model_pointer`

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- `block_pointer`

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
tagular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
```
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.25  optpp_newton

• Keywords Area
• method
• optpp_newton

Newton method based optimization

Topics

This keyword is related to the topics:

• package_optpp
• local_optimization_methods

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td>search_method</td>
<td></td>
<td>Select a search method for Newton-based optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>merit_function</td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>steplength_to_boundary</td>
<td>Controls how close to the boundary of the feasible region the algorithm is allowed to move</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>centering_parameter</td>
<td>Controls how closely the algorithm should follow the &quot;central path&quot;</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_step</td>
<td>Max change in design point</td>
<td></td>
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<tr>
<td>Optional</td>
<td>gradient_tolerance</td>
<td>Stopping criteria based on L2 norm of gradient</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
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<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
<td></td>
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<tr>
<td>Optional</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
<td></td>
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<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
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<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

This is a full Newton method that expects a gradient and a Hessian. Each of the Newton-based methods are automatically bound to the appropriate OPT++ algorithm based on the user constraint specification (unconstrained, bound-constrained, or generally-constrained). In the generally-constrained case, the Newton methods use a non-linear interior-point approach to manage the constraints.

See package_optpp for info related to all optpp methods.

**Expected HDF5 Output**
If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

See Also

These keywords may also be of interest:

- optpp_cg
- optpp_g_newton
- optpp_pds
- optpp_fd_newton
- optpp_q_newton

7.2.25.1 search_method

Select a search method for Newton-based optimizers

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>Line Search Method (Group 1)</td>
<td>value_based_line_search</td>
<td>Use only function values for line search</td>
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<tr>
<td>gradient_based_line_search</td>
<td>Set the search method to use the gradient</td>
<td></td>
<td></td>
</tr>
<tr>
<td>trust_region</td>
<td>Use trust region as the globalization strategy</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.25.1.1 `value_based_line_search`

- **Keywords Area**
  - method
  - optpp_newton
  - search_method
  - `value_based_line_search`

Use only function values for line search.

Specification

**Alias**: none

**Argument(s)**: none

**Default**: `trust_region` (unconstrained), `value_based_line_search` (bound/general constraints)

Description

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_based_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

7.2.25.1.2 `gradient_based_line_search`

- **Keywords Area**
Set the search method to use the gradient

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `search_method` control is defined for all Newton-based optimizers and is used to select between `trust_region`, `gradient_based_line_search`, and `value_based_line_search` methods. The `gradient_based_line_search` option uses the line search method proposed by [63]. This option satisfies sufficient decrease and curvature conditions; whereas, `value_base_line_search` only satisfies the sufficient decrease condition. At each line search iteration, the `gradient_based_line_search` method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the `value_based_line_search` method is preferred to the `gradient_based_line_search` method. Each of these Newton methods additionally supports the `tr_pds` selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a `trust_region` search method is the default for unconstrained problems.

### 7.2.25.1.3 trust_region

- **Keywords Area**
- **method**
- **optpp_newton**
- **search_method**
- **trust_region**

Use trust region as the globalization strategy.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used...
when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.25.1.4 tr_pds

- Keywords Area
- method
- optpp_newton
- search_method
- tr_pds

Use direct search as the local search in a trust region method

**Specification**

Alias: none

Argument(s): none

**Description**

The `search_method` control is defined for all Newton-based optimizers and is used to select between trust_region, gradient_based_line_search, and value_based_line_search methods. The gradient_based_line_search option uses the line search method proposed by[63]. This option satisfies sufficient decrease and curvature conditions; whereas, value_base_line_search only satisfies the sufficient decrease condition. At each line search iteration, the gradient_based_line_search method computes the function and gradient at the trial point. Consequently, given expensive function evaluations, the value_based_line_search method is preferred to the gradient_based_line_search method. Each of these Newton methods additionally supports the tr_pds selection for unconstrained problems. This option performs a robust trust region search using pattern search techniques. Use of a line search is the default for bound-constrained and generally-constrained problems, and use of a trust_region search method is the default for unconstrained problems.

7.2.25.2 merit_function

- Keywords Area
- method
- optpp_newton
- merit_function

Balance goals of reducing objective function and satisfying constraints
A merit_function is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

7.2.25.2.1 el_bakry

- Keywords Area
- method
- optpp_newton
- merit_function
- el_bakry

El-Bakry merit function

Specification

Alias: none
Argument(s): none

Description

The "el_bakry" merit function is the L2-norm of the first order optimality conditions for the nonlinear programming problem. The cost per linesearch iteration is n+1 function evaluations. For more information, see[20].

7.2.25.2.2 argaez_tapia

- Keywords Area
- method
- optpp_newton
- merit_function

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>The merit function by Tapia and Argaez</td>
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<td></td>
<td></td>
<td>van_shanno</td>
<td>The merit function by Vanderbei and Shanno</td>
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</table>
The merit function by Tapia and Argaez

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The "argaez_tapia" merit function can be classified as a modified augmented Lagrangian function. The augmented Lagrangian is modified by adding to its penalty term a potential reduction function to handle the perturbed complementarity condition. The cost per linesearch iteration is one function evaluation. For more information, see [83].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

7.2.25.2.3  **van_shanno**

- Keywords Area
- method
- optpp_newton
- merit_function
- van_shanno

The merit function by Vanderbei and Shanno

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The "van_shanno" merit function can be classified as a penalty function for the logarithmic barrier formulation of the nonlinear programming problem. The cost per linesearch iteration is one function evaluation. For more information see [85].

If the function evaluation is expensive or noisy, set the `merit_function` to "argaez_tapia" or "van_shanno".

7.2.25.3  **steplength_to_boundary**

- Keywords Area
- method
- optpp_newton
- steplength_to_boundary

Controls how close to the boundary of the feasible region the algorithm is allowed to move
7.2 method

Specification

Alias: none
Argument(s): REAL
Default: Merit function dependent: 0.8 (el_bakry), 0.99995 (argaez_tapia), 0.95 (van_shanno)

Description

The `steplength_to_boundary` specification is a parameter (between 0 and 1) that controls how close to the boundary of the feasible region the algorithm is allowed to move. A value of 1 means that the algorithm is allowed to take steps that may reach the boundary of the feasible region. If the user wishes to maintain strict feasibility of the design parameters this value should be less than 1. Default values are .8, .99995, and .95 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.25.4 centering_parameter

- Keywords Area
- method
- optpp_newton
- centering_parameter

Controls how closely the algorithm should follow the "central path"

Specification

Alias: none
Argument(s): REAL
Default: Merit function dependent: 0.2 (el_bakry), 0.2 (argaez_tapia), 0.1 (van_shanno)

Description

The `centering_parameter` specification is a parameter (between 0 and 1) that controls how closely the algorithm should follow the "central path". See[92] for the definition of central path. The larger the value, the more closely the algorithm follows the central path, which results in small steps. A value of 0 indicates that the algorithm will take a pure Newton step. Default values are .2, .2, and .1 for the el_bakry, argaez_tapia, and van_shanno merit functions, respectively.

7.2.25.5 max_step

- Keywords Area
- method
- optpp_newton
- max_step

Max change in design point
Specification

Alias: none
Argument(s): REAL
Default: 1000.

Description

The `max_step` control specifies the maximum step that can be taken when computing a change in the current design point (e.g., limiting the Newton step computed from current gradient and Hessian information). It is equivalent to a move limit or a maximum trust region size. The `gradient_tolerance` control defines the threshold value on the L2 norm of the objective function gradient that indicates convergence to an unconstrained minimum (no active constraints). The `gradient_tolerance` control is defined for all gradient-based optimizers.

7.2.25.6 gradient_tolerance

- Keywords Area
- method
- optpp_newton
- gradient_tolerance

Stopping criteria based on L2 norm of gradient

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The `gradient_tolerance` control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The `gradient_tolerance` control is currently defined only for the OPT++ and ROL libraries.

7.2.25.7 max_iterations

- Keywords Area
- method
- optpp_newton
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods
7.2 method

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt , local_reliability: 25; global_{reliability , interval_est , evidence} / efficient_global: 25*\( n \))

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max}_{\text{iterations}} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max}_{\text{iterations}} \) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.25.8 convergence_tolerance

- Keywords Area
- method
- optpp_newton
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4
Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.25.9 speculative

- Keywords Area
- method
- optpp_newton
- speculative

Compute speculative gradients

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no speculation

Description

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based...
optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.25.10 max_function_evaluations

- Keywords Area
- method
- optpp_newton
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior
Default value is 1000.

7.2.25.11 scaling

- Keywords Area
- method
- optpp_newton
- scaling

Turn on scaling for variables, responses, and constraints

Topics
This keyword is related to the topics:
- method_independent_controls

Specification
Alias: none
Argument(s): none
Default: no scaling

Description
Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types
There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

---

**Scales**

The *_scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *_scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

---

**Examples**

The two examples below are equivalent:

```plaintext
responses
objective_functions 3
sense "maximize"
primary_scale_types = "value"
primary_scales = 1 1 100

responses
objective_functions 3
sense "maximize"
primary_scale_types = "value" "value" "value"
primary_scales = 1 1 100
```

---

**7.2.25.12 model_pointer**

- **Keywords Area**
- **method**
• optpp_newton
• model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

evironment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
7.2.26 optpp_pds

- **Keywords Area**

- **method**

- **optpp_pds**

Simplex-based derivative free optimization method

**Topics**

This keyword is related to the topics:

- **package_optpp**

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

The direct search algorithm, PDS (parallel direct search method), supports bound constraints.

The PDS method can directly exploit asynchronous evaluations; however, this capability has not yet been implemented in Dakota.

See [package_optpp](#) for info related to all optpp methods.

### See Also

These keywords may also be of interest:

- optpp_cg
- optpp_g_newton
- optpp_fd_newton
- optpp_newton
- optpp_q_newton

### 7.2.26.1 search_scheme_size

- Keywords Area
7.2 method

- method
- optpp_pds
- search_scheme_size

Number of points to be used in the direct search template

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 32

Description

The `search_scheme_size` is defined for the PDS method to specify the number of points to be used in the direct search template.

7.2.26.2 max_iterations

- Keywords Area
- method
- optpp_pds
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed `max_iterations` iterations. In cases where there are multiple nested iterations to be controlled, `max_iterations` generally controls the outer-most context.

**Default Behavior**

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.26.3 convergence_tolerance

- Keywords Area
- method
- optpp_pds
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance
7.2.26.4 max_function_evaluations

- Keywords Area
- method
- optpp_pds
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.26.5 scaling

- Keywords Area
- method
- optpp_pds
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling

   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling

   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.
Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```
responses
  objective_functions = 3
  sense = "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions = 3
  sense = "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.26.6  model_pointer

- Keywords Area
- method
- optpp_pds
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                0.1 0.2 0.6
                0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```
Third-party optimization library integration demonstration.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>variable_tolerance</td>
<td>variable_tolerance</td>
<td>Step length-based stopping criteria for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>solution_target</td>
<td>solution_target</td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>Optional</td>
<td>options_file</td>
<td>options_file</td>
<td>Filename for a YAML file that specifies Gaussian process options</td>
</tr>
</tbody>
</table>

**Description**

This is not a method that is intended for use. Rather it is part of a documented, functional example of how to integrate a third-party optimization library. It takes an initial point, generates a list of random parameter values, evaluates the function at each point in the list, and returns the point with the lowest function value.

Documented instructions on how to integrate a third-party library, working source code, and build infrastructure can be found in $dakotaSource/packages/external/demo_tpl. Additionally, integration instructions can be found in the Developers Manual.
7.2.27.1 max_function_evaluations

Number of function evaluations allowed for optimizers

Topics
This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): INTEGER
Default: 1000

Description
The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior
Default value is 1000.

7.2.27.2 max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics
This keyword is related to the topics:

• method_independent_controls
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \(\text{max\_iterations}\) iterations. In cases where there are multiple nested iterations to be controlled, \(\text{max\_iterations}\) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \max\_function\_evaluations

7.2.27.3 convergence_tolerance

- Keywords Area
- method
- demo_tpl
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The \(\text{convergence\_tolerance}\) specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a **relative convergence tolerance** for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do. Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See **nl2sol**
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.27.4 variable_tolerance

Step length-based stopping criteria for derivative-free optimizers

**Specification**

- **Alias:** none
- **Argument(s):** REAL
- **Default:** 0.01

**Description**

The `variable tolerance` keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to `asynch_pattern_search`, `coliny_cobyla`, `coliny_pattern_search`, `coliny_solis_wets`, and `meshadaptive_search`.

**Default Behavior**

The default value varies according to method as follows:

- `asynch_pattern_search`: 1.0e-2
- `coliny_cobyla`: 1.0e-4
- `coliny_pattern_search`: 1.0e-5
- `coliny_solis_wets`: 1.0e-6
Usage Tips

It is recommended that variable_tolerance be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.

Examples

Three example method input blocks appear below.

For asynch_pattern_search:

```plaintext
method
asynch_pattern_search
  contraction_factor = 0.25
  variable_tolerance = 1.e-4
  solution_target = 1.e-6
  max_function_evaluations 500
  constraint_tolerance 1.e-6
```

For coliny_pattern_search:

```plaintext
method
coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
```

For mesh_adaptive_search:

```plaintext
method
mesh_adaptive_search
  variable_tolerance = 0.01
  seed = 1234
```

7.2.27.5 solution_target

- Keywords Area
- method
- demoTpl
- solution_target

Stopping criteria based on objective function value

Specification

Alias: solution_accuracy

Argument(s): REAL

Default: no target

Description

solution_target is a termination criterion. The algorithm will terminate when the function value falls below solution_target.
7.2.27.6 options_file

- Keywords Area
- method
- demo_tpl
- options_file

Filename for a YAML file that specifies Gaussian process options

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no advanced options file

Description

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

Examples

```
# FILE: gp_options.yaml

GP Parameters:
  Sigma Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
  scaler name: standardization
  num restarts: 20
  Nugget:
    fixed nugget: 0.0
    estimate nugget: true
  Bounds:
    lower bound: 3.17e-8
    upper bound: 1.0e-2
  gp seed: 42
  standardize response: false
  Trend:
    estimate trend: true
  Options:
    max degree: 2
    reduced basis: false
    p-norm: 1.0
    scaler type: none
    regression solver type: SVD
    verbosity: 1
  kernel type: squared exponential
  Length-scale Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
    verbosity: 1
```

7.2.28 rol

- Keywords Area
Rapid Optimization Library (ROL) is a large-scale optimization package within Trilinos.

**Topics**

This keyword is related to the topics:

- local_optimization_methods

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>variable_tolerance</td>
<td></td>
<td>Step length-based stopping criteria for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>gradient_tolerance</td>
<td></td>
<td>Stopping criteria based on L2 norm of gradient</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_-tolerance</td>
<td></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td>options_file</td>
<td></td>
<td>File containing advanced ROL options</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
</tbody>
</table>
### Description

ROL is used for the solution of optimal design, optimal control and inverse problems in large-scale engineering applications.

### Usage Tips

ROL is a general gradient-based library designed to scale well to very large problem sizes. For large problem sizes (i.e. number of variables), ROL’s trust region method and conjugate gradient methods exhibit good scalability for unconstrained problems. ROL handles equality constraints natively but converts inequality constraints into equality constraints with bounded slack variables. This has might degrade convergence for problems involving large number of inequality constraints. ROL has traditionally been applied to problems with analytic gradients (and Hessians) but can be used with Dakota-provided finite-differencing approximations to the gradient of both objective and constraints. However, a user employing these is advised to use alternative optimizers such as DOT until performance of ROL improves in future releases.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
- Calibration (when calibration_terms are specified)

#### 7.2.28.1 max_iterations

- Keywords Area
- method
- rol
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

### Topics

This keyword is related to the topics:

- method_independent_controls

### Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_{reliability, interval_est, evidence} / efficient_global: 25∗n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \text{max\_function\_evaluations}

7.2.28.2 variable_tolerance

- Keywords Area
- method
- rol
- variable_tolerance

Step length-based stopping criteria for derivative-free optimizers

Specification

Alias: none

Argument(s): REAL

Default: 0.01

Description

The \( \text{variable\_tolerance} \) keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to asynch_pattern_search, coliny_cobyla, coliny_pattern_search, coliny_solis_wets, and mesh_adaptive_search.

Default Behavior

The default value varies according to method as follows:

- asynch_pattern_search: 1.0e-2
- coliny_cobyla: 1.0e-4
- coliny_pattern_search: 1.0e-5
- coliny_solis_wets: 1.0e-6
- mesh_adaptive_search: 1.0e-6

Usage Tips

It is recommended that \( \text{variable\_tolerance} \) be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.
Examples

Three example method input blocks appear below.

For `asynch_pattern_search`:

```
method
  asynch_pattern_search
    contraction_factor = 0.25
    variable_tolerance = 1.e-4
    solution_target = 1.e-6
    max_function_evaluations = 500
    constraint_tolerance = 1.e-6
```

For `coliny_pattern_search`:

```
method
  coliny_pattern_search
    initial_delta = .2
    variable_tolerance = 1.e-4
    max_iterations = 100
    solution_accuracy = 1.e-6
    seed = 1234
    max_function_evaluations = 1000
```

For `mesh_adaptive_search`:

```
method
  mesh_adaptive_search
    variable_tolerance = 0.01
    seed = 1234
```

7.2.28.3 `gradient_tolerance`

- **Keywords Area**
- **method**
- **rol**
- **gradient_tolerance**

Stopping criteria based on L2 norm of gradient

**Specification**

**Alias:** none
**Argument(s):** REAL
**Default:** 1.e-4

**Description**

The `gradient_tolerance` control defines the threshold value on the L2 norm of the objective function gradient (for unconstrained minimum or no active constraints) or the Lagrangian gradient (for constrained minimum) that indicates convergence to a stationary point. The `gradient_tolerance` control is currently defined only for the OPT++ and ROL libraries.
7.2.28.4 constraint_tolerance

• Keywords Area
• method
• rol
• constraint_tolerance

Maximum allowable constraint violation still considered feasible

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: Library default

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algorithm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

• CONMIN optimizers: 1.0e-3
• DOT constrained optimizers: 3.0e-3
• surrogate_based_local : 1.0e-4

7.2.28.5 options_file

• Keywords Area
• method
• rol
• options_file

File containing advanced ROL options
Specification

Alias: none
Argument(s): STRING
Default: no advanced options file

Description

Allow power users to override default ROL options

Default Behavior  No advanced options used.

Usage Tips  The options_file offers a final override over other options settings, which are applied with the following precedence:

1. ROL library defaults
2. Dakota hard-coded defaults
3. Settings from Dakota input file
4. Settings from the advanced options_file

7.2.28.6  scaling

• Keywords Area
• method
• rol
• scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are
performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.28.7 model_pointer

- Keywords Area
- method
- rol
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.29  asynch_pattern_search

• Keywords Area
• method
• asynch_pattern_search

Pattern search, derivative free optimization method

Topics

This keyword is related to the topics:

• package_hopspack
• global_optimization_methods
### Specification

**Alias:** coliny_apps  
**Argument(s):** none  

**Child Keywords:**

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td><code>initial_delta</code></td>
<td>Initial step size for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>contraction_factor</code></td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>variable_tolerance</code></td>
<td>Step length-based stopping criteria for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>solution_target</code></td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>synchronization</code></td>
<td>Select how Dakota schedules a batch of concurrent function evaluations in a parallel algorithm</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>merit_function</code></td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>constraint_penalty</code></td>
<td>Multiplier for the penalty function</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>smoothing_factor</code></td>
<td>Smoothing value for smoothed penalty functions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>constraint_-tolerance</code></td>
<td>Maximum allowable constraint violation still considered feasible</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>max_function_-evaluations</code></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
</tbody>
</table>
### Description

The asynchronous parallel pattern search (APPS) algorithm [38] is a fully asynchronous pattern search technique in that the search along each offset direction continues without waiting for searches along other directions to finish. Currently, APPS only supports coordinate bases with a total of $2n$ function evaluations in the pattern, and these patterns may only contract.

### Concurrency

APPS exploits parallelism through the use of Dakota’s concurrent function evaluations. The variant of the algorithm that is currently exposed, however, limits the amount of concurrency that can be exploited. In particular, APPS can leverage an evaluation concurrency level of at most twice the number of variables. More options that allow for greater evaluation concurrency may be exposed in future releases.

#### Algorithm Behavior

- **initial_delta**: the initial step length, must be positive
- **variable_tolerance**: step length used to determine convergence, must be greater than or equal to $4.4\times10^{-16}$
- **contraction_factor**: amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1

#### Merit Functions

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-base subproblems. There are several exact and smoothed exact penalty functions that can be specified with the `merit_function` control. The options are as follows:

- **merit_max**: based on $\ell_\infty$ norm
- **merit_max_smooth**: based on smoothed $\ell_\infty$ norm
- **merit1**: based on $\ell_1$ norm
- **merit1_smooth**: based on smoothed $\ell_1$ norm
- **merit2**: based on $\ell_2$ norm
- **merit2_smooth**: based on smoothed $\ell_2$ norm
- **merit2_squared**: based on $\ell_2^2$ norm

The user can also specify the following to affect the merit functions:

- **constraint_penalty**
- **smoothing_parameter**

### Method Independent Controls

The only method independent controls that are currently mapped to APPS are:

<table>
<thead>
<tr>
<th>Optional</th>
<th>scaling</th>
<th>Turn on scaling for variables, responses, and constraints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

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Note that while APPS treats the constraint tolerance separately for linear and nonlinear constraints, we apply the same value to both if the user specifies `constraint_tolerance`.

The APPS internal display level is mapped to the Dakota `output` settings as follows:

- `debug`: display final solution, all input parameters, variable and constraint info, trial points, search directions, and execution details
- `verbose`: display final solution, all input parameters, variable and constraint info, and trial points
- `normal`: display final solution, all input parameters, variable and constraint summaries, and new best points
- `quiet`: display final solution and all input parameters
- `silent`: display final solution

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions` are specified)
- Best Nonlinear Constraints
- Calibration (when `calibration_terms` are specified)

**7.2.29.1 initial_delta**

- Keywords Area
- method
- `asynch_pattern_search`
- initial_delta

Initial step size for derivative-free optimizers

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0
Description

The `initial_delta` keyword defines the size of the first search step in derivative-free optimization methods, specifically `asynch_pattern_search`, `coliny_cobyla`, `coliny_pattern_search`, `coliny_solis_wets`, and `mesh_adaptive_search`. It is applied in an absolute sense to all search directions.

Default Behavior

The default value is 1.0.

Usage Tips

It is recommended that `initial_delta` be the approximate distance from the initial point to the solution. If this distance is unknown, it is advisable to err on the side of choosing an `initial_delta` that is too large or to not specify it. Relative application of `initial_delta` is not available unless the user scales the problem accordingly.

Examples

Three example method input blocks appear below.

For `asynch_pattern_search`:

```
method
  asynch_pattern_search
  initial_delta = .5
  contraction_factor = 0.25
  merit_function merit1_smooth
  smoothing_factor = 1.0
  constraint_tolerance = 1.e-6
```

For `coliny_pattern_search`:

```
method
  coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
```

For `mesh_adaptive_search`:

```
method
  mesh_adaptive_search
  initial_delta = 2.0
  seed = 1234
```

7.2.29.2 contraction_factor

- Keywords Area
- method
  - `asynch_pattern_search`
  - `contraction_factor`

Amount by which step length is rescaled

Specification

Alias: none
**Argument(s):** REAL  
**Default:** 0.5

**Description**

For pattern search methods, `contraction_factor` specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is $1/\text{contraction\_factor}$

7.2.29.3 variable_tolerance

- Keywords Area
- method
- asynch_pattern_search
- variable_tolerance

Step length-based stopping criteria for derivative-free optimizers

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.01

**Description**

The `variable_tolerance` keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to `asynch_pattern_search`, `coliny_cobyla`, `coliny_pattern_search`, `coliny_solis_wets`, and `mesh_adaptive_search`.

**Default Behavior**

The default value varies according to method as follows:

- `asynch_pattern_search`: $1.0\times10^{-2}$
- `coliny_cobyla`: $1.0\times10^{-4}$
- `coliny_pattern_search`: $1.0\times10^{-5}$
- `coliny_solis_wets`: $1.0\times10^{-6}$
- `mesh_adaptive_search`: $1.0\times10^{-6}$

**Usage Tips**

It is recommended that `variable_tolerance` be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.
Examples

Three example method input blocks appear below.

For `asynch_pattern_search`:

```plaintext
method
    asynch_pattern_search
    contraction_factor = 0.25
    variable_tolerance = 1.e-4
    solution_target = 1.e-6
    max_function_evaluations 500
    constraint_tolerance 1.e-6
```

For `coliny_pattern_search`:

```plaintext
method
    coliny_pattern_search
    initial_delta = .2
    variable_tolerance = 1.e-4
    max_iterations 100
    solution_accuracy = 1.e-6
    seed = 1234
    max_function_evaluations = 1000
```

For `mesh_adaptive_search`:

```plaintext
method
    mesh_adaptive_search
    variable_tolerance = 0.01
    seed = 1234
```

7.2.29.4 `solution_target`

- **Keywords Area**
- **method**
- `asynch_pattern_search`
- `solution_target`

Stopping criteria based on objective function value

**Specification**

**Alias:** `solution_accuracy`

**Argument(s):** REAL

**Default:** no target

**Description**

`solution_target` is a termination criterion. The algorithm will terminate when the function value falls below `solution_target`.

7.2.29.5 `synchronization`

- **Keywords Area**
- **method**
• `asynch_pattern_search`
• `synchronization`

Select how Dakota schedules a batch of concurrent function evaluations in a parallel algorithm.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** method specific

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Synchronization (Group 1)</th>
<th>Dakota Keyword</th>
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<tr>
<td>Required (Choose One)</td>
<td></td>
<td>blocking</td>
<td>Complete concurrent evaluation of all evaluations within a batch</td>
</tr>
<tr>
<td>nonblocking</td>
<td></td>
<td>nonblocking</td>
<td>A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm</td>
</tr>
</tbody>
</table>

**Description**

The synchronization specification can be used to specify the use of either blocking or nonblocking schedulers, where the former must block until all evaluations are completed and the latter does not block, returning only the partial set that has been completed for a particular call to the scheduler.

7.2.29.5.1 blocking

• `Keywords Area`
• `method`
• `asynch_pattern_search`
• `synchronization`
• `blocking`

Complete concurrent evaluation of all evaluations within a batch

**Specification**

**Alias:** none

**Argument(s):** none
Description

In the **blocking** case, all of the concurrent evaluations in the batch must be completed and all results are returned together to the algorithm.

These runs are reproducible, assuming use of the same seed in cases with random number generation.

### 7.2.29.5.2 nonblocking

- **Keywords Area**
- **method**
- **asynch_pattern_search**
- **synchronization**
- **nonblocking**

A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

In the **nonblocking** case, the set of running evaluations is queried for completion and a partial set of results is returned to an asynchronous algorithm, which can then generate new evaluations based on these partial results.

In all cases, the algorithm steps will be subject to parallel timing variabilities / race conditions such that these runs will not generally be repeatable regardless of seed specification.

### 7.2.29.6 merit_function

- **Keywords Area**
- **method**
- **asynch_pattern_search**
- **merit_function**

Balance goals of reducing objective function and satisfying constraints.

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: merit2_squared

**Child Keywords:**
<table>
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<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
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<td>Nonsmoothed merit function</td>
</tr>
<tr>
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<td></td>
<td>merit_max_smooth</td>
<td>Smoothed merit function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>merit1</td>
<td>Nonsmoothed merit function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>merit1_smooth</td>
<td>Smoothed merit function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>merit2</td>
<td>Nonsmoothed merit function</td>
</tr>
<tr>
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<td>merit2_smooth</td>
<td>Smoothed merit function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>merit2_squared</td>
<td>Nonsmoothed merit function</td>
</tr>
</tbody>
</table>

### Description

A **merit_function** is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

#### 7.2.29.6.1 merit_max

- **Keywords Area**
- **method**
- **asynch_pattern_search**
- **merit_function**
- **merit_max**

Nonsmoothed merit function

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-base subproblems. There are several exact and smoothed exact penalty functions.

**merit_max:** based on $\ell_\infty$ norm

#### 7.2.29.6.2 merit_max_smooth

- **Keywords Area**
- **method**
- **asynch_pattern_search**
- **merit_function**
- **merit_max_smooth**

Smoothed merit function
7.2 method

Specification

Alias: none
Argument(s): none

Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-base subproblems. There are several exact and smoothed exact penalty functions.

merit_max_smooth: based on smoothed $\ell_\infty$ norm

7.2.29.6.3 merit1

- Keywords Area
- method
- asynch_pattern_search
- merit_function
- merit1

Nonsmoothed merit function

Specification

Alias: none
Argument(s): none

Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-base subproblems. There are several exact and smoothed exact penalty functions.

- merit1: based on $\ell_1$ norm

7.2.29.6.4 merit1_smooth

- Keywords Area
- method
- asynch_pattern_search
- merit_function
- merit1_smooth

Smoothed merit function

Specification

Alias: none
Argument(s): none
APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-based subproblems. There are several exact and smoothed exact penalty functions.

7.2.29.6.5 merit1_smooth

- Keywords Area
- method
- merit1
- merit1_smooth: based on smoothed $\ell_1$ norm

7.2.29.6.6 merit2

- Keywords Area
- method
- asynch_pattern_search
- merit_function
- merit2
- merit2: based on $\ell_2$ norm

Nonsmoothed merit function

Specification

Alias: none
Argument(s): none

Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-based subproblems. There are several exact and smoothed exact penalty functions.

7.2.29.6.6 merit2_smooth

- Keywords Area
- method
- asynch_pattern_search
- merit_function
- merit2_smooth
- merit2_smooth: based on smoothed $\ell_2$ norm

Smoothed merit function

Specification

Alias: none
Argument(s): none

Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-based subproblems. There are several exact and smoothed exact penalty functions.

merit2_smooth: based on smoothed $\ell_2$ norm
7.2 method

7.2.29.6.7 merit2_squared

• Keywords Area

• method

• asynch_pattern_search

• merit_function

• merit2_squared

Nonsmoothed merit function

Specification

Alias: none
Argument(s): none

Description

APPS solves nonlinearly constrained problems by solving a sequence of linearly constrained merit function-base subproblems. There are several exact and smoothed exact penalty functions.

merit2_squared: based on ℓ_2 norm

7.2.29.7 constraint_penalty

• Keywords Area

• method

• asynch_pattern_search

• constraint_penalty

Multiplier for the penalty function

Specification

Alias: none
Argument(s): REAL
Default: 1.0

Description

Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds constraint_penalty times the sum of squares of the constraint violations to the objective function. The default value of constraint_penalty is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.
7.2.29.8 smoothing_factor

- Keywords Area
- method
- asynch_pattern_search
- smoothing_factor

Smoothing value for smoothed penalty functions

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0.0

**Description**

- smoothing_parameter: initial smoothing value for smoothed penalty functions, must be between 0 and 1 (inclusive)

7.2.29.9 constraint_tolerance

- Keywords Area
- method
- asynch_pattern_search
- constraint_tolerance

Maximum allowable constraint violation still considered feasible

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** Library default
7.2 method

Description

Sets the maximum allowable value of infeasibility that any constraint in an optimization problem may possess and
still be considered to be satisfied.

If a constraint's violation is greater than this value then it is considered to be violated by the optimization algo-
ритhm. This specification gives some control over how tightly the constraints will be satisfied at convergence of the
algorithm. However, if the value is set too small the algorithm may terminate with one or more constraints being violated.

Defaults

Defaults vary depending on the method.

- CONMIN optimizers: 1.0e-3
- DOT constrained optimizers: 3.0e-3
- surrogate_based_local : 1.0e-4

7.2.29.10 max_function_evaluations

- Keywords Area
- method
- asynch_pattern_search
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached
any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations
evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before
max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.
7.2.29.11 scaling

- Keywords Area
- method
- asynch_pattern_search
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.
2. **auto** - automatic scaling

First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

- two-sided bounds scaled into the interval [0,1];
- one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
- no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling

First, any characteristic values from the optional *scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

### Examples

The two examples below are equivalent:

```plaintext
responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value"
    primary_scales = 1 1 100

responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value" "value" "value"
    primary_scales = 1 1 100
```

7.2.29.12 **model_pointer**

- **Keywords Area**
- **method**
- **asynch_pattern_search**
- **model_pointer**

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```bash
environment
  tabular_data
    method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
        0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'II'
```
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ‘x1’ ‘x2’

interface
id_interface = ‘I1’
system async evaluation_concurrency = 5
analysis_driver = ‘text_book’

responses
response_functions = 3
no_gradients
no_hessians

7.2.30  mesh_adaptive_search

• Keywords Area
• method
• mesh_adaptive_search

Finds optimal variable values using adaptive mesh-based search

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>initial_delta</td>
<td>Initial step size for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>variable_tolerance</td>
<td>Step length-based stopping criteria for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>function_precision</td>
<td>Specify the maximum precision of the analysis code responses</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>history_file</td>
<td>Name of file where mesh adaptive search records all evaluation points.</td>
</tr>
</tbody>
</table>
Optional | display_format | Information to be reported from mesh adaptive search's internal records.

Optional | variable_-neighborhood_-search | Percentage of evaluations to do to escape local minima.

Optional | neighbor_order | Number of dimensions in which to perturb categorical variables.

Optional | display_all_-evaluations | Shows mesh adaptive search's internally held list of all evaluations

Optional | use_surrogate | Surrogate model usage mode for mesh adaptive search

Optional | max_iterations | Number of iterations allowed for optimizers and adaptive UQ methods

Optional | max_function_-evaluations | Number of function evaluations allowed for optimizers

Optional | scaling | Turn on scaling for variables, responses, and constraints

Optional | model_pointer | Identifier for model block to be used by a method

Description

The mesh adaptive direct search algorithm[8] is a derivative-free generalized pattern search in which the set of points evaluated becomes increasingly dense, leading to good convergence properties. It can handle unconstrained problems as well as those with bound constraints and general nonlinear constraints. Furthermore, it can handle continuous, discrete, and categorical variables.

Default Behavior

By default, mesh_adaptive_search operates on design variables. The types of variables can be expanded through the use of the active keyword in the variables block in the Dakota input file. Categorical variables, however, must be limited to design variables.
Expected Outputs

The best objective function value achieved and associated parameter and constraint values can be found at the end of the Dakota output. The method’s internally summarized iteration history will appear in the screen output by default, with the option to control the method’s output through Dakota’s output level. It also generates a history file containing a list of all function evaluations done.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

Additional Discussion

The mesh adaptive direct search method is made available in Dakota through the NOMAD software[2], available to the public under the GNU LGPL from http://www.gerad.ca/nomad.

Examples

The following is an example of a Dakota input file that makes use of mesh_adaptive_search to optimize the textbook function.

```plaintext
method, mesh_adaptive_search seed = 1234
variables, continuous_design = 3
    initial_point -1.0 1.5 2.0
    upper_bounds 10.0 10.0 10.0
    lower_bounds -10.0 -10.0 -10.0
    descriptors 'x1' 'x2' 'x3'
interface, direct
    analysis_driver = 'text_book'
responses, objective_functions = 1
    no_gradients
    no_hessians
```

The best function value and associated parameters are found at the end of the Dakota output.

```
<<<<<< Function evaluation summary: 674 total (674 new, 0 duplicate)
<<<<<< Best parameters = 1.0000000000e+00 1.0000000000e+00 1.0000000000e+00 1.0000000000e+00 1.0000000000e+00
<<<<<< Best objective function = 1.0735377280e-52
<<<<<< Best data captured at function evaluation 658
```

A NOMAD-generated iteration summary is also printed to the screen.
BBE OBJ

1  17.0625000000
2  1.0625000000
13  0.0625000000
24  0.0002441406
41  0.0000314713
43  0.0000028610
54  0.0000000307
83  0.0000000000
105  0.0000000000
112  0.0000000000
114  0.0000000000
135  0.0000000000
142  0.0000000000
153  0.0000000000
159  0.0000000000
171  0.0000000000
193  0.0000000000
200  0.0000000000
207  0.0000000000
223  0.0000000000
229  0.0000000000
250  0.0000000000
266  0.0000000000
282  0.0000000000
288  0.0000000000
314  0.0000000000
320  0.0000000000
321  0.0000000000
327  0.0000000000
354  0.0000000000
361  0.0000000000
372  0.0000000000
373  0.0000000000
389  0.0000000000
400  0.0000000000
417  0.0000000000
444  0.0000000000
459  0.0000000000
461  0.0000000000
488  0.0000000000
492  0.0000000000
494  0.0000000000
501  0.0000000000
518  0.0000000000
530  0.0000000000
537  0.0000000000
564  0.0000000000
566  0.0000000000
583  0.0000000000
590  0.0000000000
592  0.0000000000
604  0.0000000000
606  0.0000000000
629  0.0000000000
636  0.0000000000
658  0.0000000000
674  0.0000000000

} end of run (mesh size reached NOMAD precision)

blackbox evaluations : 674
best feasible solution : ( 1 1 1 ) h=0 f=1.073537728e-52

7.2.30.1 initial_delta

- Keywords Area
- method
- mesh_adaptive_search
7.2 method

- initial_delta

Initial step size for derivative-free optimizers

Specification

Alias: none
Argument(s): REAL
Default: 1.0

Description

The initial_delta keyword defines the size of the first search step in derivative-free optimization methods, specifically asynch_pattern_search, coliny_cobyla, coliny_pattern_search, coliny_solis_wets, and mesh_adaptive_search. It is applied in an absolute sense to all search directions.

Default Behavior

The default value is 1.0.

Usage Tips

It is recommended that initial_delta be the approximate distance from the initial point to the solution. If this distance is unknown, it is advisable to err on the side of choosing an initial_delta that is too large or to not specify it. Relative application of initial_delta is not available unless the user scales the problem accordingly.

Examples

Three example method input blocks appear below.

For asynch_pattern_search:

method
  asynch_pattern_search
      initial_delta = .5
      contraction_factor = 0.25
      merit_function merit1_smooth
      smoothing_factor = 1.0
      constraint_tolerance = 1.e-6

For coliny_pattern_search:

method
  coliny_pattern_search
      initial_delta = .2
      variable_tolerance = 1.e-4
      max_iterations 100
      solution_accuracy = 1.e-6
      seed = 1234
      max_function_evaluations = 1000

For mesh_adaptive_search

method
  mesh_adaptive_search
      initial_delta = 2.0
      seed = 1234

7.2.30.2 variable_tolerance

- Keywords Area
• method
• mesh_adaptive_search
• variable_tolerance

Step length-based stopping criteria for derivative-free optimizers

Specification

Alias: none
Argument(s): REAL
Default: 1.0e-6

Description

The variable_tolerance keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to asynch_pattern_search, coliny_cobyla, coliny_pattern_search, coliny_solis_wets, and mesh_adaptive_search.

Default Behavior

The default value varies according to method as follows:

• asynch_pattern_search: 1.0e-2
• coliny_cobyla: 1.0e-4
• coliny_pattern_search: 1.0e-5
• coliny_solis_wets: 1.0e-6
• mesh_adaptive_search: 1.0e-6

Usage Tips

It is recommended that variable_tolerance be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.

Examples

Three example method input blocks appear below.

For asynch_pattern_search:

method
  asynch_pattern_search
  contraction_factor = 0.25
  variable_tolerance = 1.e-4
  solution_target = 1.e-6
  max_function_evaluations 500
  constraint_tolerance 1.e-6

For coliny_pattern_search:

method
  coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
For `mesh_adaptive_search`

```plaintext
method
  mesh_adaptive_search
    variable_tolerance = 0.01
    seed = 1234
```

### 7.2.30.3 `function_precision`

- **Keywords Area**
- **method**
- **mesh_adaptive_search**
- **function_precision**

Specify the maximum precision of the analysis code responses

**Specification**

*Alias*: none

*Argument(s)*: REAL

*Default*: 1.0e-10

**Description**

The `function_precision` control provides the algorithm with an estimate of the accuracy to which the problem functions can be computed. This is used to prevent the algorithm from trying to distinguish between function values that differ by less than the inherent error in the calculation.

### 7.2.30.4 `seed`

- **Keywords Area**
- **method**
- **mesh_adaptive_search**
- **seed**

Seed of the random number generator

**Specification**

*Alias*: none

*Argument(s)*: INTEGER

*Default*: system-generated (non-repeatable)
The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.30.5  history_file

- Keywords Area
- method
- mesh_adaptive_search
- history_file

Name of file where mesh adaptive search records all evaluation points.

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** mads_history

**Description**

The history_file is used to specify the name of a file to which mesh adaptive direct search will write its own list of evaluated points.

**Default Behavior**

By default, mesh adaptive direct search will write the list of evaluation points in a file named mads_history.xxxx, where xxxx corresponds to a randomly generated number. Dakota's output level controls the method's level of output to file.
Examples

The example below shows the syntax for specifying the name of the history file.

```plaintext
method
  mesh_adaptive_search
  history_file = 'output.log'
  seed = 1234
```

7.2.30.6 display_format

- Keywords Area
- method
- mesh_adaptive_search
- display_format

Information to be reported from mesh adaptive search's internal records.

Specification

**Alias:** none

**Argument(s):** STRING

Description

The `display_format` keyword is used to specify the set of information to be reported by the mesh adaptive direct search method. This is information mostly internal to the method and not reported via Dakota output.

**Default Behavior**

By default, only the number of function evaluations (bbe) and the objective function value (obj) are reported. The full list of options is as follows. Note that case does not matter.

- BBE: Blackbox evaluations.
- BBO: Blackbox outputs.
- EVAL: Evaluations (includes cache hits).
- MESH_INDEX: Mesh index.
- MESH_SIZE: Mesh size parameter.
- OBJ: Objective function value.
- POLL_SIZE: Poll size parameter.
- SOL: Solution, with format iSOLj where i and j are two (optional) strings: i will be displayed before each coordinate, and j after each coordinate (except the last).
- STAT_AVG: The AVG statistic.
- STAT_SUM: The SUM statistic defined by argument.
- TIME: Wall-clock time.
- VARI: Value of variable i. The index 0 corresponds to the first variable.
Expected Outputs
A list of the requested information will be printed to the screen.

Usage Tips
This will most likely only be useful for power users who want to understand and/or report more detailed information on method behavior.

Examples
The following example shows the syntax for specifying display_format. Note that all desired information options should be listed within a single string.

```plaintext
method
  mesh_adaptive_search
  display_format 'bbe obj poll_size'
  seed = 1234
```

Below is the output reported for the above example.

```
MADS run {
  BBE OBJ POLL_SIZE
  1 17.0625000000 2.0000000000 2.0000000000 2.0000000000
  2 1.0625000000 2.0000000000 2.0000000000 2.0000000000
  13 0.0625000000 1.0000000000 1.0000000000 1.0000000000
  24 0.0000244106 0.5000000000 0.5000000000 0.5000000000
  41 0.0000314713 0.1250000000 0.1250000000 0.1250000000
  43 0.0000000037 0.1250000000 0.1250000000 0.1250000000
  54 0.0000000000 0.0781250000 0.0781250000 0.0781250000
  105 0.0000000000 0.009765625 0.009765625 0.009765625
  112 0.0000000000 0.009765625 0.009765625 0.009765625
  114 0.0000000000 0.0019531250 0.0019531250 0.0019531250
  135 0.0000000000 0.0048828128 0.0048828128 0.0048828128
  142 0.0000000000 0.0048828128 0.0048828128 0.0048828128
  153 0.0000000000 0.0048828128 0.0048828128 0.0048828128
  159 0.0000000000 0.009765625 0.009765625 0.009765625
  171 0.0000000000 0.0048828128 0.0048828128 0.0048828128
  193 0.0000000000 0.00610352 0.00610352 0.00610352
  200 0.0000000000 0.00610352 0.00610352 0.00610352
  207 0.0000000000 0.00610352 0.00610352 0.00610352
  223 0.0000000000 0.00305176 0.00305176 0.00305176
  229 0.0000000000 0.00610352 0.00610352 0.00610352
  250 0.0000000000 0.000152588 0.000152588 0.000152588
  266 0.0000000000 0.000076294 0.000076294 0.000076294
  282 0.0000000000 0.000038147 0.000038147 0.000038147
  288 0.0000000000 0.000076294 0.000076294 0.000076294
  314 0.0000000000 0.00009537 0.00009537 0.00009537
  320 0.0000000000 0.00019073 0.00019073 0.00019073
  321 0.0000000000 0.00038147 0.00038147 0.00038147
  327 0.0000000000 0.000076294 0.000076294 0.000076294
  354 0.0000000000 0.00004768 0.00004768 0.00004768
  361 0.0000000000 0.00004768 0.00004768 0.00004768
  372 0.0000000000 0.00004768 0.00004768 0.00004768
  373 0.0000000000 0.00009537 0.00009537 0.00009537
  389 0.0000000000 0.00004768 0.00004768 0.00004768
  400 0.0000000000 0.00004768 0.00004768 0.00004768
  417 0.0000000000 0.00001192 0.00001192 0.00001192
  444 0.0000000000 0.000000037 0.000000037 0.000000037
  461 0.0000000000 0.000000075 0.000000075 0.000000075
  488 0.0000000000 0.000000005 0.000000005 0.000000005
  492 0.0000000000 0.000000009 0.000000009 0.000000009
  494 0.0000000000 0.000000019 0.000000019 0.000000019
  501 0.0000000000 0.000000019 0.000000019 0.000000019
  518 0.0000000000 0.000000005 0.000000005 0.000000005
}

Dakota Version 6.15 Reference Manual generated on November 5, 2021
blackbox evaluations : 674
best feasible solution : (1 1 1) h=0 f=1.073537728e-52

See Also

These keywords may also be of interest:

- display_all_evaluations

7.2.30.7 variable_neighborhood_search

- Keywords Area
- method
- mesh_adaptive_search
- variable_neighborhood_search

Percentage of evaluations to do to escape local minima.

Specification

Alias: none
Argument(s): REAL
Default: 0.0

Description

The variable_neighborhood_search keyword is used to set the percentage (in decimal form) of function evaluations used to escape local minima. The mesh adaptive direct search method will try to perform a maximum of that percentage of the function evaluations within this more extensive search.

Default Behavior

By default, variable_neighborhood_search is not used.

Usage Tips

Using variable_neighborhood_search results in an increased number of function evaluations. If the desired result is a local minimum, the added cost is of little or no value, so the recommendation is not to use it. If the desired result is the best local minimum possible within a computational budget, then there is value in setting this parameter. Note that the higher the value, the greater the computational cost. The value should be no greater than 1.0.
Examples

The following example shows the syntax used to set variable_neighborhood_search.

```plaintext
method
  mesh_adaptive_search
    seed = 1234
    variable_neighborhood_search = 0.1
```

7.2.30.8 neighbor_order

- **Keywords Area**
- **method**
- **mesh_adaptive_search**
- **neighbor_order**

Number of dimensions in which to perturb categorical variables.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

The `neighbor_order` keyword allows the user to specify the number of categorical dimensions to perturb when determining neighboring points that will be used by the mesh adaptive direct search method to augment its search. When greater than 1, the neighbors are defined from the tensor product of the admissible 1-dimensional perturbations.

**Default Behavior**

By default, the categorical neighbors will be defined by perturbing only one categorical variable at a time (according to the corresponding adjacency_matrix; see `adjacency_matrix`) while leaving the others fixed at their current values.

**Usage Tips**

The maximum meaningful value `neighbor_order` can take on is the number of categorical variables.

**Examples**

In this example, suppose we have the following categorical variables and associated adjacency matrices.

```plaintext
variables
discrete_design_set
  real = 2
categorical 'yes' 'yes'
  num_set_values = 3 5
  set_values = 1.2 2.3 3.4 4.4 5.5 7.7
  adjacency_matrix = 1 1 0
                    1 1 1
                    0 1 1
                    1 0 1 0 1
                    1 0 1 0 1
                    1 0 1 0 1
```
Also suppose that we have the following method specification.

```plaintext
method
  mesh_adaptive_search
  seed = 1234
```

If the mesh adaptive direct search is at the point (1.2, 1.2), then the neighbors will be defined by the default 1-dimensional perturbations and would be the following:

- (2.3, 1.2)
- (1.2, 4.4)
- (1.2, 7.7)

If, instead, the method specification is the following:

```plaintext
method
  mesh_adaptive_search
  seed = 1234
  neighbor_order = 2
```

The neighbors will be defined by 2-dimensional perturbations defined from the tensor product of the 1-dimensional perturbation and would be the following:

- (2.3, 1.2)
- (2.3, 4.4)
- (2.3, 7.7)
- (1.2, 4.4)
- (1.2, 7.7)

See Also

These keywords may also be of interest:

- adjacency_matrix

7.2.30.9  display_all_evaluations

- Keywords Area
- method
- mesh_adaptive_search
- display_all_evaluations

Shows mesh adaptive search's internally held list of all evaluations

Specification

Alias: none
Argument(s): none
Default: false
Description

If set, `display_all_evaluations` will instruct the mesh adaptive direct search method to print out its own record of all evaluations. The information reported may be controlled using `display_format`.

Default Behavior

By default, mesh adaptive direct search does not report information on all evaluations, only on those for which an improvement in the objective function is found.

Expected Outputs

The information specified by `display_format` will be reported to the screen for every function evaluation.

Usage Tips

This will most likely only be useful for power users who want to understand and/or report more detailed information on method behavior.

Examples

The following example shows the syntax for specifying `display_all_evaluations`.

```plaintext
method
   mesh_adaptive_search
   display_all_evaluations
   max_function_evaluations=20
   seed = 1234

Note that the output below reports information (default for `display_format`) for all function evaluations.

MADS run {
   BBE OBJ
   1   17.0625000000
   2   1.0625000000
   3   1297.0625000000
   4   257.0625000000
   5   81.0625000000
   6   151.0625000000
   7   1051.0625000000
   8   40.0625000000
   9   17.0625000000
  10   40.0625000000
  11   1.0625000000
  12   102.0625000000
  13   0.0625000000
  14   231.0625000000
  15   16.0625000000
  16   5.0625000000
  17   16.0625000000
  18   71.0625000000
  19   0.0625000000
  20   1.0625000000
} end of run (max number of blackbox evaluations)

blackbox evaluations : 20
best feasible solution : (1 0.5 1) h=0 f=0.0625

That is in contrast with what would be reported by default.

MADS run {
   BBE OBJ
```
7.2 7.2.30.10 use_surrogate

See Also

These keywords may also be of interest:

- display_format

Surrogate model usage mode for mesh adaptive search

Specification

Alias: none

Argument(s): none

Default: optimize

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Surrogate Purpose (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>inform_search Surrogate informs evaluation order in mesh adaptive search</td>
<td>optimize</td>
<td>Surrogate is used in lieu of true model for mesh adaptive search</td>
</tr>
</tbody>
</table>

Description

The use_surrogate keyword is used to define how a surrogate model (if one is provided) is to be used by mesh_adaptive_search. There are two approaches available: inform_search uses the surrogate to sort list of trial points and subsequently the true function is evaluated on the most promising points first. Both true function and surrogate are used interchangeably within the method. optimize forces the use of a surrogate in lieu of the true model and thus the true function is never invoked except to construct the surrogate.
Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

Default Behavior

By default, `mesh_adaptive_search` follows behaviour provided by `optimize` option.

Examples

The following example shows the syntax used to set `use_surrogate`.

```
method,
  mesh_adaptive_search
  model_pointer = 'SURROGATE'
  use_surrogate inform_search

model,
  id_model = 'SURROGATE'
  surrogate global
  polynomial quadratic
  dace_method_pointer = 'SAMPLING'

variables,
  continuous_design = 3
  initial_point -1.0 1.5 2.0
  upper_bounds 10.0 10.0 10.0
  lower_bounds -10.0 -10.0 -10.0
  descriptors 'x1' 'x2' 'x3'
  discrete_design_range = 2
  initial_point 2 2
  lower_bounds 1 1
  upper_bounds 4 9
  descriptors 'y1' 'y2'
  discrete_design_set
     real = 2
     elements_per_variable = 4 5
     elements = 1.2 2.3 3.4 4.5 1.2 3.3 4.4 5.5 7.7
     descriptors 'y3' 'y4'
     integer = 2
     elements_per_variable = 2 2
     elements = 4 7 8 9
     descriptors 'z1' 'z2'

method,
  id_method = 'SAMPLING'
  model_pointer = 'TRUTH'
  sampling
    samples = 55

model,
  id_model = 'TRUTH'
  single
    interface_pointer = 'TRUE_FN'

interface,
  id_interface = 'TRUE_FN'
    direct
    analysis_driver = 'text_book'

responses,
  objective_functions = 1
  no_gradients
  no_hessians
```

7.2.30.10.1 inform_search

- Keywords Area
Surrogate informs evaluation order in mesh adaptive search

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

When `inform_search` is specified with `use_surrogate`, `mesh_adaptive_search` uses the surrogate to sort list of trial points and subsequently the true function is evaluated on the most promising points first. Both true function and surrogate are used interchangeably within the method.

**Default Behavior**

`inform_search` is not the default surrogate usage mode.

**Expected Output**

The user can expect to see both the number of true model evaluations and the number of approximation (i.e., surrogate) evaluations reported in the Dakota screen output. The former captures the sum of truth evaluations done for the surrogate construction and for the optimization.

**Usage Tips**

When `inform_search` is specified, the `maximum_function_evaluations` keyword applies to only the optimization method and does not account for evaluations needed to construct the surrogate. If the user has a strict evaluation budget, they should set `maximum_function_evaluations` such that evaluation budget = number of evaluations to construct surrogate + `maximum_function_evaluations`.

**Examples**

The following example shows the syntax used to set `use_surrogate` to optimize.

```plaintext
method,
    mesh_adaptive_search
    model_pointer = 'SURROGATE'
    use_surrogate inform_search

model,
    id_model = 'SURROGATE'
    surrogate global
    polynomial quadratic
    dace_method_pointer = 'SAMPLING'

variables,
    continuous_design = 3
    initial_point -1.0 1.5 2.0
    upper_bounds 10.0 10.0 10.0
    lower_bounds -10.0 -10.0 -10.0
    descriptors 'x1' 'x2' 'x3'
    discrete_design_range = 2
    initial_point 2 2
    lower_bounds 1 1
    upper_bounds 4 9
```
The following will appear toward the end of the screen output when Dakota is run on this example. The number of true function evaluations includes the 55 evaluations that were done to construct the surrogate (as specified in the SAMPLING method block) plus the number of truth evaluations done by `mesh_adaptive_search`.

<<<<<< Function evaluation summary (APPROX_INTERFACE): 1660 total (1660 new, 0 duplicate)
<<<<<< Function evaluation summary (TRUE_FN): 795 total (795 new, 0 duplicate)

7.2.30.10.2 optimize

- Keywords Area
- method
- mesh_adaptive_search
- use_surrogate
- optimize

Surrogate is used in lieu of true model for mesh adaptive search

Specification

Alias: none
Argument(s): none

Description

When `optimize` is specified with `use_surrogate`, `mesh_adaptive_search` will use the surrogate in lieu of the true model and thus the true function is never invoked except to construct the surrogate.

Default Behavior
**7.2 Method**

*optimize* is the default surrogate usage mode.

**Expected Output**

The user can expect to see both the number of true model evaluations and the number of approximation (i.e., surrogate) evaluations reported in the Dakota screen output. The former should equal the number of truth evaluations done for the surrogate construction.

**Examples**

The following example shows the syntax used to set *use_surrogate* to *optimize*.

```plaintext
method,
   mesh_adaptive_search
   model_pointer = 'SURROGATE'
   use_surrogate optimize

model,
   id_model = 'SURROGATE'
   surrogate global
   polynomial quadratic
   dace_method_pointer = 'SAMPLING'

variables,
   continuous_design = 3
      initial_point -1.0  1.5  2.0
      upper_bounds  10.0 10.0 10.0
      lower_bounds -10.0 -10.0 -10.0
      descriptors  'x1'  'x2'  'x3'
   discrete_design_range = 2
      initial_point  2   2   2
      lower_bounds  1   1   1
      upper_bounds  4   9   9
      descriptors   'y1'  'y2'
   discrete_design_set
      real = 2
         elements_per_variable = 4 5
         elements = 1.2 2.3 3.4 4.5 1.2 3.3 4.4 5.5 7.7
         descriptors  'y3'  'y4'
      integer = 2
         elements_per_variable = 2 2
         elements = 4 7 8 9
         descriptors  'z1'  'z2'

method,
   id_method = 'SAMPLING'
   model_pointer = 'TRUTH'
   sampling
      samples = 55

model,
   id_model = 'TRUTH'
   single
      interface_pointer = 'TRUE_FN'

interface,
   id_interface = 'TRUE_FN'
      direct
      analysis_driver = 'text_book'

responses,
   objective_functions = 1
   no_gradients
   no_hessians
```

The following will appear toward the end of the screen output when Dakota is run on this example. The number of true function evaluations includes only the 55 evaluations that were done to construct the surrogate (as specified in the SAMPLING method block).

```
<<<<<< Function evaluation summary (APPROX_INTERFACE): 221 total (221 new, 0 duplicate)
```
7.2.30.11   max_iterations

-  Keywords Area
-  method
-  mesh_adaptive_search
-  max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

-  method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

-  max_function_evaluations

7.2.30.12   max_function_evaluations

-  Keywords Area
-  method
-  mesh_adaptive_search
-  max_function_evaluations

Number of function evaluations allowed for optimizers
7.2 method

Topics
This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description
The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior
Default value is 1000.

7.2.30.13 scaling

• Keywords Area
• method
• mesh_adaptive_search
• scaling

Turn on scaling for variables, responses, and constraints

Topics
This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description
Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

### Scale Types

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval $[0,1]$;
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.
   
   Logarithmic scaling is not available for linear constraints.
   
   When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The `*scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `*scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto, log` - optional
- `value` - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10\cdot\text{DBL_MIN}$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses  
  objective_functions 3  
  sense "maximize"  
  primary_scale_types = "value"  
  primary_scales = 1 1 100

responses  
  objective_functions 3  
  sense "maximize"  
  primary_scale_types = "value" "value" "value"  
  primary_scales = 1 1 100
```

7.2.30.14 model_pointer

- Keywords Area
- method
- mesh_adaptive_search
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.
Examples

environment
  tabular_data
  method_pointer = 'UQ'
method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
  sample_type lhs
  distribution cumulative
model
  id_model = 'SURR'
    surrogate global,
      dace_method_pointer = 'DACE'
      polynomial quadratic
method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2
model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'
variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'
interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.31 nowpac

  • Keywords Area
  • method
  • nowpac

Gradient-free inequality-constrained optimization using Nonlinear Optimization With Path Augmented Constraints (NOWPAC).

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

NOWPAC is a provably-convergent gradient-free optimization method from MIT that solves a series of trust region surrogate-based subproblems to generate improving steps. Due to its use of an interior penalty scheme and enforcement of strict feasibility, it does not support linear or nonlinear equality constraints. As opposed to the stochastic version (SNOWPAC), NOWPAC does not currently support a feasibility restoration mode, so it is necessary to start from a feasible design.

Note: (S)NOWPAC is not configured with Dakota by default and requires a separate installation of the NOWPAC distribution from MIT, combined with its TPLs of Eigen and NLOPT.

### Examples

```dakota
method
nowpac
  max_function_evaluations = 1000
  convergence_tolerance = 1e-4
  trust_region
    initial_size = 0.10
    minimum_size = 1.0e-6
    contract_threshold = 0.25
    expand_threshold = 0.75
    contraction_factor = 0.50
    expansion_factor = 1.50
```

### 7.2.31.1 trust_region

- Keywords Area
- method
- nowpac
- trust_region
Use trust region as the globalization strategy.

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>initial_size</td>
<td>initial_size</td>
<td>Trust region initial size (relative to bounds)</td>
</tr>
<tr>
<td>Optional</td>
<td>minimum_size</td>
<td>minimum_size</td>
<td>Trust region minimum size</td>
</tr>
<tr>
<td>Optional</td>
<td>contract_threshold</td>
<td>contract_threshold</td>
<td>Shrink trust region if trust region ratio is below this value</td>
</tr>
<tr>
<td>Optional</td>
<td>expand_threshold</td>
<td>expand_threshold</td>
<td>Expand trust region if trust region ratio is above this value</td>
</tr>
<tr>
<td>Optional</td>
<td>contraction_factor</td>
<td>contraction_factor</td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td>expansion_factor</td>
<td>expansion_factor</td>
<td>Trust region expansion factor</td>
</tr>
</tbody>
</table>

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.31.1.1 initial_size

- Keywords Area
- method
- nowpac
- trust_region
- initial_size

Trust region initial size (relative to bounds)
7.2 method

Specification

**Alias:** none

**Argument(s):** REALLIST

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

All of these specifications are REAL scalars, with the exception of the specification of `initial_size`, which is a REALLIST. This array corresponds to the case when there are more than 2 model forms or discretizations within a model hierarchy. The default `initial_size` involves a recursive halving of the global bounds for each trust region in the hierarchy: e.g., a scalar value of .5 for a single trust region managing two model forms/discretizations, or an array of (.125, .25, .5) for four model forms/discretizations (three trust regions ordered from the lowest to highest fidelity surrogate, with model four as truth).

7.2.31.1.2 minimum_size

- **Keywords** Area
  - method
  - nowpac
  - trust_region
  - minimum_size

Trust region minimum size

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-6

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The
command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.31.1.3 contract_threshold

- Keywords Area
- method
- nowpac
- trust_region
- contract_threshold

Shrink trust region if trust region ratio is below this value

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0.25

**Description**

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.31.1.4 expand_threshold

- Keywords Area
- method
- nowpac
- trust_region
- expand_threshold

Expand trust region if trust region ratio is above this value

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0.75
Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.31.1.5 contraction_factor

- Keywords Area
- method
- nowpac
- trust_region
- contraction_factor

Amount by which step length is rescaled

Specification

Alias: none

Argument(s): REAL

Default: 0.25

Description

For pattern search methods, contraction_factor specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is \( 1 / \text{contraction_factor} \)

7.2.31.1.6 expansion_factor

- Keywords Area
- method
- nowpac
- trust_region
- expansion_factor

Trust region expansion factor
Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 2.0

**Description**

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expand_threshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.31.2 max_iterations

- **Keywords Area**
- **method**
- **nowpac**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_reliability, interval_est, evidence) / efficient_global: 25+n

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed `max_iterations` iterations. In cases where there are multiple nested iterations to be controlled, `max_iterations` generally controls the outer-most context.

**Default Behavior**

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.31.3 max_function_evaluations

- Keywords Area
- method
- nowpac
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.31.4 scaling

- Keywords Area
- method
- nowpac
- scaling

Turn on scaling for variables, responses, and constraints
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `∗scales` in the variables and responses blocks. When the `scaling` keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `∗scales` keyword, which gives characteristic values (divisors)
- a `∗scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].
3. **log** - logarithmic scaling

   First, any characteristic values from the optional *scales* specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN*. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.31.5 **model_pointer**

- **Keywords Area**
- **method**
- **nowpac**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
7.2 method

```
    analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians
```

7.2.32 snowpac

- Keywords Area
- method
- snowpac

Stochastic version of NOWPAC that incorporates error estimates and noise mitigation.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>trust_region</td>
<td>Use trust region as the globalization strategy.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_function_-_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

Description

NOWPAC is a provably-convergent gradient-free optimization method from MIT that solves a series of trust region surrogate-based subproblems to generate improving steps. The stochastic version is SNOWPAC, which incorporates noise estimates in its objective and inequality constraints. SNOWPAC modifies its trust region controls and
adds smoothing from a Gaussian process surrogate in order to mitigate noise. SNOWPAC also supports a feasibility restoration mode, so it is not necessary to start from a feasible design.

Note: (S)NOWPAC is not configured with Dakota by default and requires a separate installation of the NOWPAC distribution from MIT, combined with its TPLs of Eigen and NLOPT.

Examples

Relative to the NOWPAC specification, SNOWPAC supports a seed control for repeatability of runs and also requires the return of error estimates from the underlying evaluator (e.g., UQ method such as Monte Carlo sampling).

```yaml
method,
   snowpac
   seed = 2504
   max_function_evaluations = 1000
   convergence_tolerance = 1e-4
   trust_region
      initial_size = 0.10
      minimum_size = 1.0e-6
      contract_threshold = 0.25
      expand_threshold = 0.75
      contraction_factor = 0.50
      expansion_factor = 1.50

7.2.32.1 seed
```

- Keywords Area
- method
- snowpac
- seed

Seed of the random number generator

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.
Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.32.2 trust_region

• Keywords Area
• method
• snowpac
• trust_region

Use trust region as the globalization strategy.

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>initial_size</td>
<td>initial_size</td>
<td>Trust region initial size (relative to bounds)</td>
</tr>
<tr>
<td>Optional</td>
<td>minimum_size</td>
<td>minimum_size</td>
<td>Trust region minimum size</td>
</tr>
<tr>
<td>Optional</td>
<td>contract_threshold</td>
<td>contract_threshold</td>
<td>Shrink trust region if trust region ratio is below this value</td>
</tr>
<tr>
<td>Optional</td>
<td>expand_threshold</td>
<td>expand_threshold</td>
<td>Expand trust region if trust region ratio is above this value</td>
</tr>
<tr>
<td>Optional</td>
<td>contraction_factor</td>
<td>contraction_factor</td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td>expansion_factor</td>
<td>expansion_factor</td>
<td>Trust region expansion factor</td>
</tr>
</tbody>
</table>

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.
7.2.32.2.1  initial_size

- Keywords Area
- method
- snowpac
- trust_region
- initial_size

Trust region initial size (relative to bounds)

Specification

Alias: none
Argument(s): REALIST

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

All of these specifications are REAL scalars, with the exception of the specification of initial_size, which is a REALIST. This array corresponds to the case when there are more than 2 model forms or discretizations within a model hierarchy. The default initial_size involves a recursive halving of the global bounds for each trust region in the hierarchy; e.g., a scalar value of .5 for a single trust region managing two model forms/discretizations, or an array of (.125, .25, .5) for four model forms/discretizations (three trust regions ordered from the lowest to highest fidelity surrogate, with model four as truth).

7.2.32.2.2  minimum_size

- Keywords Area
- method
- snowpac
- trust_region
- minimum_size

Trust region minimum size

Specification

Alias: none
Argument(s): REAL
Default: 1.e-6
Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.32.3 contract_threshold

- Keywords Area
  - method
  - snowpac
  - trust_region
  - contract_threshold

Shrink trust region if trust region ratio is below this value

Specification

Alias: none

Argument(s): REAL

Default: 0.25

Description

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.32.4 expand_threshold

- Keywords Area
  - method
  - snowpac
  - trust_region
  - expand_threshold
Expand trust region if trust region ratio is above this value

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.75

**Description**

The trust_region optional group specification can be used to specify the initial size of the trust region (using initial_size) relative to the total variable bounds, the minimum size of the trust region (using minimum_size), the contraction factor for the trust region size (using contraction_factor) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using expansion_factor) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using contract_threshold) and the trust region size expansion threshold (using expand_threshold). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The command contract_threshold sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The command expand_threshold determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.32.2.5 contraction_factor

- Keywords Area  
- method  
- snowpac  
- trust_region  
- contraction_factor

Amount by which step length is rescaled

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.25

**Description**

For pattern search methods, contraction_factor specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is $1/\text{contraction\_factor}$

7.2.32.2.6 expansion_factor

- Keywords Area  
- method  
- snowpac
7.2 method

- trust_region
- expansion_factor

Trust region expansion factor

Specification

Alias: none
Argument(s): REAL
Default: 2.0

Description

The `trust_region` optional group specification can be used to specify the initial size of the trust region (using `initial_size`) relative to the total variable bounds, the minimum size of the trust region (using `minimum_size`), the contraction factor for the trust region size (using `contraction_factor`) used when the surrogate model is performing poorly, and the expansion factor for the trust region size (using `expansion_factor`) used when the surrogate model is performing well. Two additional commands are the trust region size contraction threshold (using `contract_threshold`) and the trust region size expansion threshold (using `expandThreshold`). These two commands are related to what is called the trust region ratio, which is the actual decrease in the truth model divided by the predicted decrease in the truth model in the current trust region. The `contract_threshold` sets the minimum acceptable value for the trust region ratio, i.e., values below this threshold cause the trust region to shrink for the next SBL iteration. The `expand_threshold` determines the trust region value above which the trust region will expand for the next SBL iteration.

7.2.32.3 max_iterations

- Keywords Area
- method
- snowpac
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global{(reliability, interval_est, evidence)} / efficient_global: 25×n)
Description
The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed $\text{max\_iterations}$ iterations. In cases where there are multiple nested iterations to be controlled, $\text{max\_iterations}$ generally controls the outer-most context.

Default Behavior
Default value is 100.

See Also
These keywords may also be of interest:

- $\text{max\_function\_evaluations}$

7.2.32.4 $\text{max\_function\_evaluations}$

- Keywords Area
- method
- snowpac
- $\text{max\_function\_evaluations}$

Number of function evaluations allowed for optimizers

Topics
This keyword is related to the topics:

- $\text{method\_independent\_controls}$

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description
The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed $\text{max\_function\_evaluations}$ evaluations. See also $\text{max\_iterations}$.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before $\text{max\_function\_evaluations}$ was exceeded.

Default Behavior
Default value is 1000.
7.2 method

7.2.32.5 scaling

- Keywords Area
- method
- snowpac
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value·characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.
2. **auto** - automatic scaling

First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

- two-sided bounds scaled into the interval \([0,1]\);
- one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
- no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into \([0,1]\).

3. **log** - logarithmic scaling

First, any characteristic values from the optional \(*_\text{scales}\) specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The \(*_\text{scales}\) keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding \(*\text{scale_TYPES}\), as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0e10\cdot\text{DBL_MIN}\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

### Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.32.6 **model_pointer**

- **Keywords Area**
- **method**
- **snowpac**
- **model_pointer**

Identifier for model block to be used by a method
 Topics
This keyword is related to the topics:

- block_pointer

 Specification

 Alias: none

 Argument(s): STRING

 Default: method use of last model parsed (or use of default model if none parsed)

 Description

 The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

 Default Behavior

 If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

 Usage Tips

 When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

 See block_pointer for details about pointers.

 Examples

 environment
  tabular_data
  method_pointer = 'UQ'

 method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
  samples = 10
  seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
                  0.1 0.2 0.6
                  0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

 model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

 method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

 model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
variables
  uniform_uncertain = 2
  lower_bounds = 0.0.
  upper_bounds = 1.1.
  descriptors = ‘x1’ ‘x2’

interface
  id_interface = ’I1’
  system asynch evaluation_concurrency = 5
  analysis_driver = ’text_book’

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.33 moga

  • Keywords Area
  • method
  • moga

Multi-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)

Topics

This keyword is related to the topics:

  • package_jega

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Select the fitness type for JEGA methods</td>
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<td>niching_type</td>
<td></td>
<td>Specify the type of niching pressure</td>
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<tr>
<td>Optional</td>
<td>convergence_type</td>
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<td>Optional</td>
<td>postprocessor_type</td>
<td></td>
<td>Post process the final solution from moga</td>
</tr>
<tr>
<td>Parameter</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
<td></td>
</tr>
<tr>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
<td></td>
</tr>
<tr>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
<td></td>
</tr>
<tr>
<td>population_size</td>
<td>Set the initial population size in JEGA methods</td>
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<td></td>
</tr>
<tr>
<td>log_file</td>
<td>Specify the name of a log file</td>
<td></td>
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</tr>
<tr>
<td>print_each_pop</td>
<td>Print every population to a population file</td>
<td></td>
<td></td>
</tr>
<tr>
<td>initialization_type</td>
<td>Specify how to initialize the population</td>
<td></td>
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<td>crossover_type</td>
<td>Select a crossover type for JEGA methods</td>
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<td>mutation_type</td>
<td>Select a mutation type for JEGA methods</td>
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</tr>
<tr>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
<td></td>
</tr>
<tr>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
<td></td>
<td></td>
</tr>
<tr>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
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</tr>
</tbody>
</table>

### Description

*moga* stands for Multi-objective Genetic Algorithm, which is a global optimization method that does Pareto optimization for multiple objectives. It supports general constraints and a mixture of real and discrete variables.

### Constraints

*moga* can utilize linear constraints using the keywords: *
- linear_inequality_constraint_matrix
- linear_inequality_lower_bounds
- linear_inequality_upper_bounds
- linear_inequality_scale_types
- linear_inequality_scales
- linear_equality_constraint_matrix
- linear_equality_targets
- linear_equality_scale_types
- linear_equality_scales

### Configuration
The genetic algorithm configurations are:

1. fitness
2. replacement
3. niching
4. convergence
5. postprocessor
6. initialization
7. crossover
8. mutation
9. population size

The steps followed by the algorithm are listed below. The configurations will effect how the algorithm completes each step.

**Stopping Criteria**

The *moga* method respects the `max_iterations` and `max_function_evaluations` method independent controls to provide integer limits for the maximum number of generations and function evaluations, respectively. The algorithm also stops when convergence is reached. This involves repeated assessment of the algorithm’s progress in solving the problem, until some criterion is met.

The specification for convergence in a moga can either be `metric_tracker` or can be omitted all together. If omitted, no convergence algorithm will be used and the algorithm will rely on stopping criteria only.

**Expected Outputs**

The *moga* method respects the `output` method independent control to vary the amount of information presented to the user during execution.

The final results are written to the Dakota tabular output. Additional information is also available - see the `log-file` and `print_each_pop` keywords.

Note that moga and SOGA create additional output files during execution. "finaldata.dat" is a file that holds the final set of Pareto optimal solutions after any post-processing is complete. "discards.dat" holds solutions that were discarded from the population during the course of evolution.

It can often be useful to plot objective function values from these files to visually see the Pareto front and ensure that finaldata.dat solutions dominate discards.dat solutions. The solutions are written to these output files in the format "Input1...InputN..Output1...OutputM".

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions` are specified)
- Best Nonlinear Constraints
- Calibration (when `calibration_terms` are specified)

**Important Notes**

The pool of potential members is the current population and the current set of offspring.

Choice of fitness assessors is strongly related to the type of replacement algorithm being used and can have a profound effect on the solutions selected for the next generation.
If using the fitness types `layer_rank` or `domination_count`, it is strongly recommended that you use the replacement_type `below_limit` (although the roulette wheel selectors can also be used).

The functionality of the domination_count selector of JEGA v1.0 can now be achieved using the domination_count fitness type and below_limit replacement type.

**Theory**

The basic steps of the moga algorithm are as follows:

1. Initialize the population
2. Evaluate the population (calculate the values of the objective function and constraints for each population member)
3. Loop until converged, or stopping criteria reached
   (a) Perform crossover
   (b) Perform mutation
   (c) Evaluate the new population
   (d) Assess the fitness of each member in the population
   (e) Replace the population with members selected to continue in the next generation
   (f) Apply niche pressure to the population
   (g) Test for convergence
4. Perform post processing

If moga is used in a hybrid optimization method (which requires one optimal solution from each individual optimization method to be passed to the subsequent optimization method as its starting point), the solution in the Pareto set closest to the "utopia" point is given as the best solution. This solution is also reported in the Dakota output.

This "best" solution in the Pareto set has minimum distance from the utopia point. The utopia point is defined as the point of extreme (best) values for each objective function. For example, if the Pareto front is bounded by (1,100) and (90,2), then (1,2) is the utopia point. There will be a point in the Pareto set that has minimum L2-norm distance to this point, for example (10,10) may be such a point.

If moga is used in a method which may require passing multiple solutions to the next level (such as the surrogate_based_global method or hybrid methods), the orthogonal_distance postprocessor type may be used to specify the distances between each solution value to winnow down the solutions in the full Pareto front to a subset which will be passed to the next iteration.

**See Also**

These keywords may also be of interest:

- soga

**7.2.33.1 fitness_type**

- Keywords Area
- method
- moga
- fitness_type

Select the fitness type for JEGA methods
Specification

Alias: none

Argument(s): none

Default: domination_count

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Fitness Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>layer_rank</td>
<td>Assign each member to a layer, based on domination the rank based on layers</td>
</tr>
<tr>
<td></td>
<td></td>
<td>domination_count</td>
<td>Rank each member by the number of members that dominate it</td>
</tr>
</tbody>
</table>

Description

The two JEGA methods use different fitness types, which are described on their respective pages.

7.2.33.1.1 layer_rank

- Keywords Area
- method
- moga
- fitness_type
- layer_rank

Assign each member to a layer, based on domination the rank based on layers

Specification

Alias: none

Argument(s): none

Description

The fitness_type: layer_rank has been specifically designed to avoid problems with aggregating and scaling objective function values and transforming them into a single objective.

The layer_rank fitness assessor works by assigning all non-dominated designs a layer of 0, then from what remains, assigning all the non-dominated a layer of 1, and so on until all designs have been assigned a layer. The values are negated to follow the higher-is-better fitness convention.

Use of the below_limit selector with the layer_rank fitness assessor has the effect of keeping all those designs whose layer is below a certain threshold again subject to the shrinkage limit.
7.2.33.1.2 domination_count

- Keywords Area
- method
- moga
- fitness_type
- domination_count

Rank each member by the number of members that dominate it

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `fitness_type: domination_count` has been specifically designed to avoid problems with aggregating and scaling objective function values and transforming them into a single objective.

Instead, the `domination_count` fitness assessor works by ordering population members by the negative of the number of designs that dominate them. The values are negated in keeping with the convention that higher fitness is better.

The `layer_rank` fitness assessor works by assigning all non-dominated designs a layer of 0, then from what remains, assigning all the non-dominated a layer of 1, and so on until all designs have been assigned a layer. Again, the values are negated for the higher-is-better fitness convention.

Use of the `below_limit` selector with the `domination_count` fitness assessor has the effect of keeping all designs that are dominated by fewer than a limiting number of other designs subject to the shrinkage limit.

Using it with the `layer_rank` fitness assessor has the effect of keeping all those designs whose layer is below a certain threshold again subject to the shrinkage limit.

7.2.33.2 replacement_type

- Keywords Area
- method
- moga
- replacement_type

Select a replacement type for JEGA methods

**Specification**

**Alias:** none  
**Argument(s):** none

**Default:** below_limit

**Child Keywords:**
Required/Optional | Description of Group | Dakota Keyword | Dakota Keyword Description
--- | --- | --- | ---
Required (Choose One) | Replacement Type (Group 1) | elitist | Use the best designs to form a new population

| Replacement Type (Group 1) | roulette_wheel | Replace population |
| unique_roulette_wheel | Replace population |
| below_limit | Limit number of designs dominating those kept |

**Description**

Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The *replacement_type* of *roulette_wheel* or *unique_roulette_wheel* may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the *layer_rank* and *domination_count*, the recommended selector is the *below_limit* selector. The *below_limit* replacement will only keep designs that are dominated by fewer than a limiting number of other designs.

In *roulette_wheel* replacement, each design is conceptually allotted a portion of a wheel proportional to its fitness relative to the fitnesses of the other Designs. Then, portions of the wheel are chosen at random and the design occupying those portions are duplicated into the next population. Those Designs allotted larger portions of the wheel are more likely to be selected (potentially many times). *unique_roulette_wheel* replacement is the same as *roulette_wheel* replacement, with the exception that a design may only be selected once. The *below_limit* selector attempts to keep all designs for which the negated fitness is below a certain limit. The values are negated to keep with the convention that higher fitness is better. The inputs to the *below_limit* selector are the limit as a real value, and a *shrinkage_percentage* as a real value. The *shrinkage_percentage* defines the minimum amount of selections that will take place if enough designs are available. It is interpreted as a percentage of the population size that must go on to the subsequent generation. To enforce this, *below_limit* makes all the selections it would make anyway and if that is not enough, it takes the remaining that it needs from the best of what is left (effectively raising its limit as far as it must to get the minimum number of selections). It continues until it has made enough selections. The *shrinkage_percentage* is designed to prevent extreme decreases in the population size at any given generation, and thus prevent a big loss of genetic diversity in a very short time. Without a shrinkage limit, a small group of “super” designs may appear and quickly pull the population down to a size on the order of the limiting value. In this case, all the diversity of the population is lost and it is expensive to re-diversify and spread the population. The

The *replacement_type* for a SOGA may be *roulette_wheel*, *unique_roulette_wheel*, *elitist*, or *favor_feasible*. The *elitist* selector simply chooses the required number of designs taking the most fit. For example, if 100 selections are requested, then the top 100 designs as ranked by fitness will be selected and the remaining will be discarded. The *favor_feasible* replacement type first considers feasibility as a selection criteria. If that does not produce a “winner” then it moves on to considering fitness value. Because of this, any fitness assessor used with the *favor_feasible* selector must only account objectives in the creation of fitness. Therefore, there is such a fitness assessor and it’s use is enforced when the *favor_feasible* selector is chosen. In that case, and if the output level is set high enough, a message will be presented indicating that the *weighted_sum_only* fitness assessor will be used.

7.2.33.2.1 *elitist*

- **Keywords Area**
- **method**
- **moga**
- **replacement_type**
- **elitist**
Use the best designs to form a new population

Specification

Alias: none
Argument(s): none

Description

The elitist (default) setting creates a new population using (a) the replacement_size best individuals from the current population, (b) and population_size - replacement_size individuals randomly selected from the newly generated individuals. It is possible in this case to lose a good solution from the newly generated individuals if it is not randomly selected for replacement; however, the default new_solutions_generated value is set such that the entire set of newly generated individuals will be selected for replacement.

7.2.33.2.2 roulette_wheel

- Keywords Area
- method
- moga
- replacement_type
- roulette_wheel

Replace population

Specification

Alias: none
Argument(s): none

Description

Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The replacement_type of roulette_wheel or unique_roulette_wheel may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the layer_rank and domination_count, the recommended selector is the below_limit selector. The below_limit replacement will only keep designs that are dominated by fewer than a limiting number of other designs. The replacement_type of favor_feasible is specific to a SOGA. This replacement operator will always prefer a more feasible design to a less feasible one. Beyond that, it favors solutions based on an assigned fitness value which must have been installed by the weighted sum only fitness assessor (see the discussion below).

7.2.33.2.3 unique_roulette_wheel

- Keywords Area
- method
- moga
- replacement_type
- unique_roulette_wheel
Replace population

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The `replacement_type` of `roulette_wheel` or `unique_roulette_wheel` may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the `layer_rank` and `domination_count`, the recommended selector is the `below_limit` selector. The `below_limit` replacement will only keep designs that are dominated by fewer than a limiting number of other designs. The `replacement_type` of `favor_feasible` is specific to a SOGA. This replacement operator will always prefer a more feasible design to a less feasible one. Beyond that, it favors solutions based on an assigned fitness value which must have been installed by the weighted sum only fitness assessor (see the discussion below).

### 7.2.33.2.4 `below_limit`

- **Keywords Area**
- **method**
- **moga**
- **replacement_type**
- **below_limit**

Limit number of designs dominating those kept

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 6

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>shrinkage_fraction</td>
<td>Decrease the population size by a percentage</td>
</tr>
</tbody>
</table>

**Description**

The `below_limit` replacement will only keep designs that are dominated by fewer than a limiting number of other designs.

### 7.2.33.2.4.1 `shrinkage_fraction`

- **Keywords Area**
- **method**
7.2 method

- moga
- replacement_type
- below_limit
- shrinkage_fraction

Decrease the population size by a percentage

Specification

Alias: shrinkage_percentage
Argument(s): REAL
Default: 0.9

Description

As of JEGA v2.0, all replacement types are common to both MOGA and SOGA. They include the roulette-wheel, unique_roulette_wheel, elitist, and below_limit selectors. In roulette_wheel replacement, each design is conceptually allotted a portion of a wheel proportional to its fitness relative to the fitnesses of the other Designs. Then, portions of the wheel are chosen at random and the design occupying those portions are duplicated into the next population. Those Designs allotted larger portions of the wheel are more likely to be selected (potentially many times). unique_roulette_wheel replacement is the same as roulette_wheel replacement, with the exception that a design may only be selected once. The below_limit selector attempts to keep all designs for which the negated fitness is below a certain limit. The values are negated to keep with the convention that higher fitness is better. The inputs to the below_limit selector are the limit as a real value, and a shrinkage_percentage as a real value. The shrinkage_percentage defines the minimum amount of selections that will take place if enough designs are available. It is interpreted as a percentage of the population size that must go on to the subsequent generation. To enforce this, below_limit makes all the selections it would make anyway and if that is not enough, it takes the remaining that it needs from the best of what is left (effectively raising its limit as far as it must to get the minimum number of selections). It continues until it has made enough selections. The shrinkage_percentage is designed to prevent extreme decreases in the population size at any given generation, and thus prevent a big loss of genetic diversity in a very short time. Without a shrinkage limit, a small group of "super" designs may appear and quickly cull the population down to a size on the order of the limiting value. In this case, all the diversity of the population is lost and it is expensive to re-diversify and spread the population. The elitist selector simply chooses the required number of designs taking the most fit. For example, if 100 selections are requested, then the top 100 designs as ranked by fitness will be selected and the remaining will be discarded.

7.2.33 niching_type

- Keywords Area
- method
- moga
- niching_type

Specify the type of niching pressure
Specification

Alias: none

Argument(s): none

Default: No niche pressure

Child Keywords:

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<td>radial</td>
<td>Set niching distance to percentage of non-dominated range</td>
</tr>
<tr>
<td></td>
<td>max_designs</td>
<td></td>
<td>Enforce minimum Euclidean distance between designs</td>
</tr>
</tbody>
</table>

Description

The purpose of niching is to encourage differentiation along the Pareto frontier and thus a more even and uniform sampling.

This is typically accomplished by discouraging clustering of design points in the performance space. In JEGA, the application of niche pressure occurs as a secondary selection operation. The niching operator is given a chance to perform a pre-selection operation prior to the operation of the selection (replacement) operator, and is then called to perform niching on the set of designs that were selected by the selection operator.

The radial nicher takes information input from the user to compute a minimum allowable distance between designs in the performance space and acts as a secondary selection operator whereby it enforces this minimum distance. The distance nicher requires that solutions must be separated from other solutions by a minimum distance in each dimension (vs. Euclidean distance for the radial niching). After niching is complete, all designs in the population will be at least the minimum distance from one another in all directions.

The radial niche pressure applicator works by enforcing a minimum Euclidean distance between designs in the performance space at each generation. The algorithm proceeds by starting at the (or one of the) extreme designs along objective dimension 0 and marching through the population removing all designs that are too close to the current design. One exception to the rule is that the algorithm will never remove an extreme design which is defined as a design that is maximal or minimal in all but 1 objective dimension (for a classical 2 objective problem, the extreme designs are those at the tips of the non-dominated frontier). The distance nicher enforces a minimum distance in each dimension.

The designs that are removed by the nicher are not discarded. They are buffered and re-inserted into the population during the next pre-selection operation. This way, the selector is still the only operator that discards designs and the algorithm will not waste time “re-filling” gaps created by the nicher.

The radial nicher requires as input a vector of fractions with length equal to the number of objectives. The elements of the vector are interpreted as percentages of the non-dominated range for each objective defining a minimum distance to all other designs. All values should be in the range (0, 1). The minimum allowable distance between any two designs in the performance space is the Euclidian (simple square-root-sum-of-squares calculation) distance defined by these percentages. The distance nicher has a similar input vector requirement, only the distance is the minimum distance in each dimension.

The max_designs niche pressure applicator is designed to choose a limited number of solutions to remain in the population. That number is specified by num_designs. It does so in order to balance the tendency for populations to grow very large and thus consuming too many computer resources. It operates by ranking designs according to their fitness standing and a computed count of how many other designs are too close to them.
close is a function of the supplied niche_vector, which specifies the minimum distance between any two points in
the performance space along each dimension individually. Once the designs are all ranked, the top c\num_designs
designs are kept in the population and the remaining ones are buffered or discarded. Note that like other niching
operators, this one will not discard an extreme design.

7.2.33.3.1 radial

• Keywords Area
  • method
  • moga
  • niching_type
  • radial

Set niching distance to percentage of non-dominated range

Specification

Alias: none
Argument(s): REALLIST
Default: 0.01 for all objectives

Description

The radial nicher requires as input a vector of fractions with length equal to the number of objectives. The
elements of the vector are interpreted as percentages of the non-dominated range for each objective defining a
minimum distance to all other designs. All values should be in the range (0, 1). The minimum allowable distance
between any two designs in the performance space is the Euclidian (simple square-root-sum-of-squares calculation)
distance defined by these percentages. The distance nicher has a similar input vector requirement, only the
distance is the minimum distance in each dimension.

7.2.33.3.2 distance

• Keywords Area
  • method
  • moga
  • niching_type
  • distance

Enforce minimum Euclidean distance between designs

Specification

Alias: none
Argument(s): REALLIST
Currently, the only niche pressure operators available are the radial nicher, the distance nicher, and the max_designs nicher. The radial niche pressure applicator works by enforcing a minimum Euclidean distance between designs in the performance space at each generation. The algorithm proceeds by starting at the (or one of the) extreme designs along objective dimension 0 and marching through the population removing all designs that are too close to the current design. One exception to the rule is that the algorithm will never remove an extreme design which is defined as a design that is maximal or minimal in all but 1 objective dimension (for a classical 2 objective problem, the extreme designs are those at the tips of the non-dominated frontier). The distance nicher enforces a minimum distance in each dimension.

The designs that are removed by the nicher are not discarded. They are buffered and re-inserted into the population during the next pre-selection operation. This way, the selector is still the only operator that discards designs and the algorithm will not waste time “re-filling” gaps created by the nicher.

### max_designs

**Limit number of solutions to remain in the population**

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>num_designs</td>
<td></td>
<td>num_designs</td>
<td>Limit the number of solutions</td>
</tr>
</tbody>
</table>

**Description**

The max_designs niche pressure applicator is designed to choose a limited number of solutions to remain in the population. That number is specified by num_designs. It does so in order to balance the tendency for populations to grow very large and thus consuming too many computer resources. It operates by ranking designs according to their fitness standing and a computed count of how many other designs are too close to them. Too close is a function of the supplied niche_vector, which specifies the minimum distance between any two points in the performance space along each dimension individually. Once the designs are all ranked, the top c \ num_designs designs are kept in the population and the remaining ones are buffered or discarded. Note that like other niching operators, this one will not discard an extreme design.

### num_designs

**Limit number of solutions to remain in the population**

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
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<th>Dakota Keyword Description</th>
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<tr>
<td>num_designs</td>
<td></td>
<td>num_designs</td>
<td>Limit the number of solutions</td>
</tr>
</tbody>
</table>

**Description**

The max_designs niche pressure applicator is designed to choose a limited number of solutions to remain in the population. That number is specified by num_designs. It does so in order to balance the tendency for populations to grow very large and thus consuming too many computer resources. It operates by ranking designs according to their fitness standing and a computed count of how many other designs are too close to them. Too close is a function of the supplied niche_vector, which specifies the minimum distance between any two points in the performance space along each dimension individually. Once the designs are all ranked, the top c \ num_designs designs are kept in the population and the remaining ones are buffered or discarded. Note that like other niching operators, this one will not discard an extreme design.
• max_designs
• num_designs

Limit the number of solutions

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

The `max_designs` niche pressure applicator is designed to choose a limited number of solutions to remain in the population. That number is specified by `num_designs`. It does so in order to balance the tendency for populations to grow very large and thus consuming too many computer resources. It operates by ranking designs according to their fitness standing and a computed count of how many other designs are too close to them. Too close is a function of the supplied `niche_vector`, which specifies the minimum distance between any two points in the performance space along each dimension individually. Once the designs are all ranked, the top `\ num_designs` designs are kept in the population and the remaining ones are buffered or discarded. Note that like other niching operators, this one will not discard an extreme design.

7.2.33.4 convergence_type

• Keywords Area
  • method
  • moga
  • convergence_type

Select the convergence type for JEGA methods

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** average_fitness_tracker

**Child Keywords:**

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<td>Track changes in the non-dominated frontier</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>percent_change</td>
<td>Define the convergence criterion for JEGA methods</td>
</tr>
</tbody>
</table>
### Description

The two JEGA methods use different convergence types, which are described on their respective pages. All the convergence types are modified by the optional keywords `percent_change` and `num_generations`.

#### 7.2.33.4.1 metric_tracker

- **Keywords Area**
- **method**
- **moga**
- **convergence_type**
- **metric_tracker**

Track changes in the non-dominated frontier

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** `metric_tracker`

### Description

The moga converger (`metric_tracker`) operates by tracking various changes in the non-dominated frontier from generation to generation. When the changes occurring over a user specified number of generations fall below a user specified threshold, the algorithm stops.

If `metric_tracker` is specified, then a `percent_change` and `num_generations` must be supplied as well. These are listed as optional keywords in the input spec.

### Theory

The `metric_tracker` converger tracks 3 metrics specific to the non-dominated frontier from generation to generation. All 3 of these metrics are computed as percent changes between the generations. In order to compute these metrics, the converger stores a duplicate of the non-dominated frontier at each generation for comparison to the non-dominated frontier of the next generation.

The first metric is one that indicates how the expanse of the frontier is changing. The expanse along a given objective is defined by the range of values existing within the non-dominated set. The expansion metric is computed by tracking the extremes of the non-dominated frontier from one generation to the next. Any movement of the extreme values is noticed and the maximum percentage movement is computed as:

\[
Em = \max_{j} \frac{\text{abs}(\text{range}(j, i) - \text{range}(j, i-1))}{\text{range}(j, i-1)} 
\]

where `Em` is the max expansion metric, `j` is the objective function index, `i` is the current generation number, and `nof` is the total number of objectives. The range is the difference between the largest value along an objective and the smallest when considering only non-dominated designs.
The second metric monitors changes in the density of the non-dominated set. The density metric is computed as the number of non-dominated points divided by the hypervolume of the non-dominated region of space. Therefore, changes in the density can be caused by changes in the number of non-dominated points or by changes in size of the non-dominated space or both. The size of the non-dominated space is computed as:

\[ V_{ps}(i) = \prod_{j=1}^{\text{range}(j, i)} j = 1, \text{nof} \]

where \( V_{ps}(i) \) is the hypervolume of the non-dominated space at generation i and all other terms have the same meanings as above.

The density of the a given non-dominated space is then:

\[ D_{ps}(i) = \frac{Pct(i)}{V_{ps}(i)} \]

where \( Pct(i) \) is the number of points on the non-dominated frontier at generation i.

The percentage increase in density of the frontier is then calculated as

\[ Cd = \frac{\text{abs}(D_{ps}(i) - D_{ps}(i-1))}{D_{ps}(i-1)} \]

where \( Cd \) is the change in density metric.

The final metric is one that monitors the “goodness” of the non-dominated frontier. This metric is computed by considering each design in the previous population and determining if it is dominated by any designs in the current population. All that are determined to be dominated are counted. The metric is the ratio of the number that are dominated to the total number that exist in the previous population.

As mentioned above, each of these metrics is a percentage. The tracker records the largest of these three at each generation. Once the recorded percentage is below the supplied percent change for the supplied number of generations consecutively, the algorithm is converged.

7.2.33.4.2 percent_change

- Keywords Area
- method
- moga
- convergence_type
- percent_change

Define the convergence criterion for JEGA methods

Specification

Alias: none

Argument(s): REAL

Default: 0.1

Description

The \( \text{percent\_change} \) is the threshold beneath which convergence is attained whereby it is compared to the metric value computed.

7.2.33.4.3 num_generations

- Keywords Area
- method
• moga

• convergence_type

• num_generations

Define the convergence criterion for JEGA methods

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 10

**Description**

The `num_generations` is the number of generations over which the metric value should be tracked. Convergence will be attained if the recorded metric is below `percent_change` for `num_generations` consecutive generations.

7.2.33.5 postprocessor_type

• Keywords Area

• method

• moga

• postprocessor_type

Post process the final solution from moga

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** No post-processing of solutions

**Child Keywords:**

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<thead>
<tr>
<th>Required/Optional Required</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>orthogonal_distance</td>
<td>Get subset of Pareto front based on distance</td>
</tr>
</tbody>
</table>

**Description**

The purpose of this operation is to perform any needed data manipulations on the final solution deemed necessary. Currently the `orthogonal_distance` is the only one. It reduces the final solution set size such that a minimum distance in each direction exists between any two designs.
7.2.33.5.1 orthogonal_distance

- Keywords Area
- method
- moga
- postprocessor_type
- orthogonal_distance

Get subset of Pareto front based on distance

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 0.01 for all objectives

**Description**

Note that MOGA and SOGA create additional output files during execution. "finaldata.dat" is a file that holds the final set of Pareto optimal solutions after any post-processing is complete. "discards.dat" holds solutions that were discarded from the population during the course of evolution. It can often be useful to plot objective function values from these files to visually see the Pareto front and ensure that finaldata.dat solutions dominate discards.dat solutions. The solutions are written to these output files in the format "Input1...InputN..Output1...OutputM". If MOGA is used in a hybrid optimization meta-iteration (which requires one optimal solution from each individual optimization method to be passed to the subsequent optimization method as its starting point), the solution in the Pareto set closest to the "utopia" point is given as the best solution. This solution is also reported in the Dakota output. This "best" solution in the Pareto set has minimum distance from the utopia point. The utopia point is defined as the point of extreme (best) values for each objective function. For example, if the Pareto front is bounded by (1,100) and (90,2), then (1,2) is the utopia point. There will be a point in the Pareto set that has minimum L2-norm distance to this point, for example (10,10) may be such a point. In SOGA, the solution that minimizes the single objective function is returned as the best solution. If moga is used in meta-iteration which may require passing multiple solutions to the next level (such as the surrogate_based_global or hybrid methods), the orthogonal_distance postprocessor type may be used to specify the distances between each solution value to winnow down the solutions in the full Pareto front to a subset which will be passed to the next iteration.

7.2.33.6 max_iterations

- Keywords Area
- method
- moga
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25∗n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.33.7 max_function_evaluations

- Keywords Area
- method
- moga
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior
Default value is 1000.

7.2.3.8 scaling

- Keywords Area
- method
- moga
- scaling

Turn on scaling for variables, responses, and constraints

Topics
This keyword is related to the topics:
- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description
Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types
There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component
   
   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional \(*_scales\) specification are applied. Then logarithm base 10 scaling is applied.
   
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

---

**Scales**

The \(*_scales\) keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding \(*_scale_types\), as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto**, **log** - optional
- **value** - required

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0\times\text{DBL_MIN}\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

---

**Examples**

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

---

7.2.33.9 **population_size**

- **Keywords Area**
- **method**
Set the initial population size in JEGA methods

Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 50  

**Description**

The number of designs in the initial population is specified by the `population_size`. Note that the `population_size` only sets the size of the initial population. The population size may vary in the JEGA methods according to the type of operators chosen for a particular optimization run.

Also note that some initializers may not strictly respect the initial population size. See the SOGA `flat_file` or MOGA `flat_file` initializer for an example.

7.2.33.10 log_file

- **Keywords Area**
- **method**
- **moga**
- **log_file**

Specify the name of a log file

Specification

**Alias:** none  
**Argument(s):** STRING  
**Default:** JEGAGlobal.log  

**Description**

New as of JEGA v2.0 is the introduction of the `log_file` specification. JEGA now uses a logging library to output messages and status to the user. JEGA can be configured at build time to log to both the console window and a text file, one or the other, or neither. The `log_file` input is a string name of a file into which to log. If the build was configured without file logging in JEGA, this input is ignored. If file logging is enabled and no `log_file` is specified, the default file name of JEGAGlobal.log is used.

7.2.33.11 print_each_pop

- **Keywords Area**
- **method**
- moga
- print_each_pop

Print every population to a population file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** No printing

**Description**

New to JEGA v2.0 is the introduction of the `print_each_pop` specification. It serves as a flag and if supplied, the population at each generation will be printed to a file named "population<GEN#>.dat" where `<GEN#>` is the number of the current generation.

7.2.33.12 `initialization_type`

- Keywords Area
- method
- moga
- initialization_type

Specify how to initialize the population

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** unique_random

**Child Keywords:**

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<tr>
<td></td>
<td></td>
<td>simple_random</td>
<td>Create random initial solutions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>unique_random</td>
<td>Create random initial solutions, but enforce uniqueness (default)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>flat_file</td>
<td>Read initial solutions from file</td>
</tr>
</tbody>
</table>

**Description**

The `initialization_type` defines how the initial population is created for the GA. There are three types:

1. `simple_random`
2. **unique_random** *(default)*

3. **flat_file**

Setting the size for the flat_file initializer has the effect of requiring a minimum number of designs to create. If this minimum number has not been created once the files are all read, the rest are created using the unique_random initializer and then the simple_random initializer if necessary.

### 7.2.33.12.1 simple_random

- **Keywords Area**
- **method**
- **moga**
  - **initialization_type**
  - **simple_random**

Create random initial solutions

#### Specification

**Alias**: none  
**Argument(s)**: none

#### Description

**simple_random** creates initial solutions with random variable values according to a uniform random number distribution. It gives no consideration to any previously generated designs.

### 7.2.33.12.2 unique_random

- **Keywords Area**
- **method**
- **moga**
  - **initialization_type**
  - **unique_random**

Create random initial solutions, but enforce uniqueness *(default)*

#### Specification

**Alias**: none  
**Argument(s)**: none

#### Description

**unique_random** is the same as **simple_random**, except that when a new solution is generated, it is checked against the rest of the solutions. If it duplicates any of them, it is rejected.
7.2.33.12.3 flat_file

- Keywords Area
- method
- moga
- initialization_type
- flat_file

Read initial solutions from file

**Specification**

**Alias:** none  
**Argument(s):** STRING

**Description**

`flat_file` allows the initial population to be read from a flat file. If `flat_file` is specified, a file name must be given.

Variables can be delimited in the flat file in any way you see fit with a few exceptions. The delimiter must be the same on any given line of input with the exception of leading and trailing whitespace. So a line could look like: 1.1, 2.2, 3.3 for example but could not look like: 1.1, 2.2, 3.3. The delimiter can vary from line to line within the file which can be useful if data from multiple sources is pasted into the same input file. The delimiter can be any string that does not contain any of the characters .+-dDeE or any of the digits 0-9. The input will be read until the end of the file. The algorithm will discard any configurations for which it was unable to retrieve at least the number of design variables. The objective and constraint entries are not required but if ALL are present, they will be recorded and the design will be tagged as evaluated so that evaluators may choose not to re-evaluate them.

Setting the size for this initializer has the effect of requiring a minimum number of designs to create. If this minimum number has not been created once the files are all read, the rest are created using the `unique_random` initializer and then the `simple_random` initializer if necessary. If more designs are found in the initialization files than specified by the initial size, then the initial size is ignored and all designs read out of the files are kept in the initial population.

7.2.33.13 crossover_type

- Keywords Area
- method
- moga
- crossover_type

Select a crossover type for JEGA methods

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** shuffle_random

**Child Keywords:**
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<tr>
<th>Required/Optional</th>
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<td>Use bit switching for crossover events</td>
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<td>multi_point_parameterized_binary</td>
<td>Use bit switching to crossover each design variable</td>
</tr>
<tr>
<td></td>
<td></td>
<td>multi_point_real</td>
<td>Perform crossover in real valued genome</td>
</tr>
<tr>
<td></td>
<td></td>
<td>shuffle_random</td>
<td>Perform crossover by choosing design variable(s)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>crossover_rate</td>
<td>Specify the probability of a crossover event</td>
</tr>
</tbody>
</table>

**Description**

There are many crossover types available. **multi_point_binary** crossover requires an integer number, \( N \), of crossover points. This crossover type performs a bit switching crossover at \( N \) crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). **multi_point_parameterized_binary** crossover is similar in that it performs a bit switching crossover routine at \( N \) crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at \( N \) locations. **multi_point_real** crossover performs a variable switching crossover routing at \( N \) crossover points in the real real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by \( 10^6 \) and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

The final crossover type is **shuffle_random**. This crossover type performs crossover by choosing design variables at random from a specified number of parents enough times that the requested number of children are produced. For example, consider the case of 3 parents producing 2 children. This operator would go through and for each design variable, select one of the parents as the donor for the child. So it creates a random shuffle of the parent design variable values. The relative numbers of children and parents are controllable to allow for as much mixing as desired. The more parents involved, the less likely that the children will wind up exact duplicates of the parents.

All crossover types take a **crossover_rate**. The crossover rate is used to calculate the number of crossover operations that take place. The number of crossovers is equal to the rate \( \times \) population_size.

**7.2.33.13.1 multi_point_binary**

- Keywords Area
- method
- moga
- crossover_type
- multi_point_binary

Use bit switching for crossover events
Specification

Alias: none
Argument(s): INTEGER

Description

There are many crossover types available. multi_point_binary crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). multi_point_parameterized_binary crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. multi_point_real crossover performs a variable switching crossover routing at N crossover points in the real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by $10^{6}$ and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

7.2.33.13.2 multi_point_parameterized_binary

- Keywords Area
- method
- moga
- crossover_type
- multi_point_parameterized_binary

Use bit switching to crossover each design variable

Specification

Alias: none
Argument(s): INTEGER

Description

There are many crossover types available. multi_point_binary crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). multi_point_parameterized_binary crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. multi_point_real crossover performs a variable switching crossover routing at N crossover points in the real real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by $10^{6}$ and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.
7.2.33.3 multi_point_real

- Keywords Area
- method
- moga
- crossover_type
- multi_point_real

Perform crossover in real valued genome

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

There are many crossover types available. `multi_point_binary` crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). `multi_point_parameterized_binary` crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. `multi_point_real` crossover performs a variable switching crossover routing at N crossover points in the real real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by $10^6$ and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

7.2.33.13.4 shuffle_random

- Keywords Area
- method
- moga
- crossover_type
- shuffle_random

Perform crossover by choosing design variable(s)

**Specification**

**Alias:** none

**Argument(s):** none

Child Keywords:
### shufflerandom-crossover

The final crossover type is `shuffle_random`. This crossover type performs crossover by choosing design variables at random from a specified number of parents enough times that the requested number of children are produced. For example, consider the case of 3 parents producing 2 children. This operator would go through and for each design variable, select one of the parents as the donor for the child. So it creates a random shuffle of the parent design variable values. The relative numbers of children and parents are controllable to allow for as much mixing as desired. The more parents involved, the less likely that the children will wind up exact duplicates of the parents.

#### num_parents

- **Keywords Area**
- **method**
- **moga**
- **crossover_type**
- **shuffle_random**
- **num_parents**

Number of parents in random shuffle crossover

**Specification**

- **Alias:** none
- **Argument(s):** INTEGER
- **Default:** 2

**Description**

Number of parents in random shuffle crossover

#### num_offspring

- **Keywords Area**
- **method**
- **moga**
- **crossover_type**
- **shuffle_random**
- **num_offspring**

Number of offspring in random shuffle crossover

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
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<tr>
<td>Optional</td>
<td>num_parents</td>
<td>num_parents</td>
<td>Number of parents in random shuffle crossover</td>
</tr>
<tr>
<td>Optional</td>
<td>num_offspring</td>
<td>num_offspring</td>
<td>Number of offspring in random shuffle crossover</td>
</tr>
</tbody>
</table>
7.2 method

Specification
Alias: none
Argument(s): INTEGER
Default: 2

Description
Number of offspring in random shuffle crossover

7.2.33.13.5 crossover_rate

- Keywords Area
- method
- moga
- crossover_type
- crossover_rate

Specify the probability of a crossover event

Specification
Alias: none
Argument(s): REAL
Default: 0.8

Description
The crossover_type controls what approach is employed for combining parent genetic information to create offspring, and the crossover_rate specifies the probability of a crossover operation being performed to generate a new offspring. The SCOLIB EA method supports three forms of crossover, two_point, blend, and uniform, which generate a new individual through combinations of two parent individuals. Two-point crossover divides each parent into three regions, where offspring are created from the combination of the middle region from one parent and the end regions from the other parent. Since the SCOLIB EA does not utilize bit representations of variable values, the crossover points only occur on coordinate boundaries, never within the bits of a particular coordinate. Uniform crossover creates offspring through random combination of coordinates from the two parents. Blend crossover generates a new individual randomly along the multidimensional vector connecting the two parents.

7.2.33.14 mutation_type

- Keywords Area
- method
- moga
- mutation_type

Select a mutation type for JEGA methods
Specification

**Alias:** none

**Argument(s):** none

**Default:** replace_uniform

### Child Keywords:

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
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<tr>
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<td>bit_random</td>
<td><strong>Mutation Type</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>replace_uniform</td>
<td>Use uniformly distributed value over range of parameter</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_normal</td>
<td>Set mutation offset to use a normal distribution</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_cauchy</td>
<td>Use a Cauchy distribution for the mutation offset</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_uniform</td>
<td>Set mutation offset to use a uniform distribution</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>mutation_rate</td>
<td>Set probability of a mutation</td>
<td></td>
</tr>
</tbody>
</table>

### Description

Five mutation types are available for selection by keyword: replace_uniform, bit_random, offset_cauchy, offset_normal, and offset_uniform. They are described in greater detail on their respective keyword pages.

The offset_∗ mutators all act by adding a random "offset" to a variable value. The random amount has a mean of zero in all cases. The size of the offset is controlled using the mutation_scale keyword, which is interpreted differently for each offset_∗ type.

The rate of mutations for all types is controlled using the mutation_rate. The rate is applied differently in each mutation_type.

#### 7.2.33.14.1 bit_random

- Keywords Area
- method
- moga
- mutation_type
- bit_random

Mutate by flipping a random bit

### Specification

**Alias:** none

**Argument(s):** none
Description

The *bit_random* mutator introduces random variation by first converting a randomly chosen variable of a randomly chosen design into a binary string. It then flips a randomly chosen bit in the string from a 1 to a 0 or visa versa. In this mutation scheme, the resulting value has more probability of being similar to the original value.

7.2.33.14.2 replace_uniform

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **replace_uniform**

Use uniformly distributed value over range of parameter

Specification

**Alias**: none

**Argument(s)**: none

Description

*replace_uniform* introduces random variation by first randomly choosing a design variable of a randomly selected design and reassigning it to a random valid value for that variable. No consideration of the current value is given when determining the new value.

7.2.33.14.3 offset_normal

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **offset_normal**

Set mutation offset to use a normal distribution

Specification

**Alias**: none

**Argument(s)**: none

Child Keywords:
Description

The offset_normal mutator introduces random variation by adding a Gaussian random amount to a variable value. The random amount has a standard deviation dependent on the mutation_scale.

7.2.33.14.3.1 mutation_scale

- Keywords Area
- method
- moga
- mutation_type
- offset_normal
- mutation_scale

Scales mutation across range of parameter

Specification

Alias: none

Argument(s): REAL

Default: 0.15

Description

The mutation_scale is a fraction in the range [0, 1] and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected mutation_type. For offset_normal and offset_cauchy, mutation_scale is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For offset_uniform, the range of possible deviation amounts is +/- 1/2 * (mutation_scale * variable range).

7.2.33.14.4 offset_cauchy

- Keywords Area
- method
- moga
- mutation_type
- offset_cauchy

Use a Cauchy distribution for the mutation offset

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

The `offset_cauchy` mutator introduces random variation by adding a Cauchy random amount to a variable value. The random amount has a standard deviation dependent on the `mutation_scale`.

#### 7.2.33.14.4.1 mutation_scale

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **offset_cauchy**
- **mutation_scale**

Scales mutation across range of parameter

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.15

**Description**

The `mutation_scale` is a fraction in the range [0, 1] and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected `mutation_type`. For `offset_normal` and `offset_cauchy`, `mutation_scale` is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For `offset_uniform`, the range of possible deviation amounts is +/- 1/2 * (mutation_scale * variable range).

#### 7.2.33.14.5 offset_uniform

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **offset_uniform**

Set mutation offset to use a uniform distribution.
Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

The **offset_uniform** mutator introduces random variation by adding a uniform random amount to a variable value. The random amount depends on the **mutation_scale**.

#### 7.2.33.14.5.1 mutation_scale

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **offset_uniform**
- **mutation_scale**

Scales mutation across range of parameter

### Specification

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.15

### Description

The **mutation_scale** is a fraction in the range \([0, 1]\) and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected **mutation_type**. For **offset_normal** and **offset_cauchy**, **mutation_scale** is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For **offset_uniform**, the range of possible deviation amounts is \(\pm 1/2 \times (\text{mutation_scale} \times \text{variable range})\).

#### 7.2.33.14.6 mutation_rate

- **Keywords Area**
- **method**
- **moga**
- **mutation_type**
- **mutation_rate**

Set probability of a mutation

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Specification

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.08

**Description**

All mutation types have a `mutation_rate`, which controls the number of mutations performed. For `replace_uniform` and all the `offset_*` types, the number of mutations performed is the product of `mutation_rate` and `population_size`. For `bit_random`, it’s the product of the `mutation_rate`, number of design variables, and population size.

7.2.33.15 seed

- **Keywords Area**  
- **method**  
- **moga**  
- **seed**

Seed of the random number generator

Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```
7.2.33.16 convergence_tolerance

- Keywords Area
- method
- moga
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.33.17 model_pointer

- Keywords Area
- method
• moga
  • model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:
  • block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
        0.1 0.2 0.6
        0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
    id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
    id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.34 soga

- Keywords Area
- method
- soga

Single-objective Genetic Algorithm (a.k.a Evolutionary Algorithm)

Topics

This keyword is related to the topics:

- package_jega
- global_optimization_methods

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>fitness_type</td>
<td>Select the fitness type for JEGA methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>replacement_type</td>
<td>Select a replacement type for JEGA methods</td>
</tr>
</tbody>
</table>
### Optional Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>convergence_type</code></td>
<td>Select the convergence type for JEGA methods</td>
</tr>
<tr>
<td><code>max_iterations</code></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td><code>max_function_evaluations</code></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td><code>scaling</code></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td><code>population_size</code></td>
<td>Set the initial population size in JEGA methods</td>
</tr>
<tr>
<td><code>log_file</code></td>
<td>Specify the name of a log file</td>
</tr>
<tr>
<td><code>print_each_pop</code></td>
<td>Print every population to a population file</td>
</tr>
<tr>
<td><code>initialization_type</code></td>
<td>Specify how to initialize the population</td>
</tr>
<tr>
<td><code>crossover_type</code></td>
<td>Select a crossover type for JEGA methods</td>
</tr>
<tr>
<td><code>mutation_type</code></td>
<td>Select a mutation type for JEGA methods</td>
</tr>
<tr>
<td><code>seed</code></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td><code>convergence_tolerance</code></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td><code>model_pointer</code></td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

### Description

`soga` stands for Single-objective Genetic Algorithm, which is a global optimization method that supports general constraints and a mixture of real and discrete variables. `soga` is part of the JEGA library.

**Constraints**  
`soga` can utilize linear constraints.

**Configuration**

The genetic algorithm configurations are:
1. fitness
2. replacement
3. convergence
4. initialization
5. crossover
6. mutation
7. population size

The pool of potential members is the current population and the current set of offspring. Choice of fitness assessors is strongly related to the type of replacement algorithm being used and can have a profound effect on the solutions selected for the next generation.

**Stopping Criteria**

The soga method respects the `max_iterations` and `max_function_evaluations` method independent controls to provide integer limits for the maximum number of generations and function evaluations, respectively.

The algorithm also stops when convergence is reached. This involves repeated assessment of the algorithm's progress in solving the problem, until some criterion is met.

**Expected Outputs**

The soga method respects the `output` method independent control to vary the amount of information presented to the user during execution.

The final results are written to the Dakota tabular output. Additional information is also available - see the `log_file` and `print_each_pop` keywords.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions`) are specified)
- Best Nonlinear Constraints
- Calibration (when `calibration_terms` are specified)

**Theory**

The basic steps of the soga algorithm are as follows:

1. Initialize the population
2. Evaluate the population (calculate the values of the objective function and constraints for each population member)
3. Loop until converged, or stopping criteria reached
   a. Perform crossover
   b. Perform mutation
   c. Evaluate the new population
   d. Assess the fitness of each member in the population
   e. Replace the population with members selected to continue in the next generation
   f. Test for convergence
Select the fitness type for JEGA methods

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** merit_function

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required</td>
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<td>merit_function</td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
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<tr>
<td>Optional</td>
<td></td>
<td>constraint_penalty</td>
<td>Multiplier for the penalty function</td>
</tr>
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</table>

**Description**

The two JEGA methods use different fitness types, which are described on their respective pages.

7.2.34.1 merit_function

- **Keywords Area**
- **method**
- **soga**
- **fitness_type**
- **merit_function**

Balance goals of reducing objective function and satisfying constraints

**Specification**

**Alias:** none

**Argument(s):** none
Description

A merit function is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.

7.2.34.1.2 constraint_penalty

- Keywords Area
- method
- soga
- fitness_type
- constraint_penalty

Multiplier for the penalty function

Specification

Alias: none
Argument(s): REAL
Default: 1.0

Description

The merit function fitness assessor uses an exterior penalty function formulation to penalize infeasible designs. The specification allows the input of a constraint_penalty which is the multiplier to use on the constraint violations.

7.2.34.2 replacement_type

- Keywords Area
- method
- soga
- replacement_type

Select a replacement type for JEGA methods

Specification

Alias: none
Argument(s): none
Default: elitist

Child Keywords:
Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The replacement_type of roulette_wheel or unique_roulette_wheel may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the layer_rank and domination_count, the recommended selector is the below_limit selector. The below_limit replacement will only keep designs that are dominated by fewer than a limiting number of other designs.

In roulette_wheel replacement, each design is conceptually allotted a portion of a wheel proportional to its fitness relative to the fitnesses of the other Designs. Then, portions of the wheel are chosen at random and the design occupying those portions are duplicated into the next population. Those Designs allotted larger portions of the wheel are more likely to be selected (potentially many times). unique_roulette_wheel replacement is the same as roulette_wheel replacement, with the exception that a design may only be selected once. The below_limit selector attempts to keep all designs for which the negated fitness is below a certain limit. The values are negated to keep with the convention that higher fitness is better. The inputs to the below_limit selector are the limit as a real value, and a shrinkage_percentage as a real value. The shrinkage_percentage defines the minimum amount of selections that will take place if enough designs are available. It is interpreted as a percentage of the population size that must go on to the subsequent generation. To enforce this, below_limit makes all the selections it would make anyway and if that is not enough, it takes the remaining that it needs from the best of what is left (effectively raising its limit as far as it must to get the minimum number of selections). It continues until it has made enough selections. The shrinkage_percentage is designed to prevent extreme decreases in the population size at any given generation, and thus prevent a big loss of genetic diversity in a very short time. Without a shrinkage limit, a small group of “super” designs may appear and quickly pull the population down to a size on the order of the limiting value. In this case, all the diversity of the population is lost and it is expensive to re-diversify and spread the population. The

The replacement_type for a SOGA may be roulette_wheel, unique_roulette_wheel, elitist, or favor_feasible. The elitist selector simply chooses the required number of designs taking the most fit. For example, if 100 selections are requested, then the top 100 designs as ranked by fitness will be selected and the remaining will be discarded. The favor_feasible replacement type first considers feasibility as a selection criteria. If that does not produce a “winner” then it moves on to considering fitness value. Because of this, any fitness assessor used with the favor_feasible selector must only account objectives in the creation of fitness. Therefore, there is such a fitness assessor and it’s use is enforced when the favor_feasible selector is chosen. In that case, and if the output level is set high enough, a message will be presented indicating that the weighted_sum_only fitness assessor will be used.

7.2.34.2.1 elitist

- Keywords Area
- method
- soga
- replacement_type
- elitist

Use the best designs to form a new population
7.2 method

Specification

Alias: none
Argument(s): none

Description

The elitist (default) setting creates a new population using (a) the replacement_size best individuals from the current population, (b) and population_size - replacement_size individuals randomly selected from the newly generated individuals. It is possible in this case to lose a good solution from the newly generated individuals if it is not randomly selected for replacement; however, the default new_solutions_generated value is set such that the entire set of newly generated individuals will be selected for replacement.

7.2.34.2.2 favor_feasible

- Keywords Area
- method
- soga
- replacement_type
- favor_feasible

Prioritize feasible designs

Specification

Alias: none
Argument(s): none

Description

This replacement operator will always prefer a more feasible design to a less feasible one. Beyond that, it favors solutions based on an assigned fitness value which must have been installed by the weighted sum only fitness assessor.

The favor_feasible replacement type first considers feasibility as a selection criteria. If that does not produce a "winner" then it moves on to considering fitness value. Because of this, any fitness assessor used with the favor_feasible selector must only account objectives in the creation of fitness. Therefore, there is such a fitness assessor and it's use is enforced when the favor_feasible selector is chosen. In that case, and if the output level is set high enough, a message will be presented indicating that the weighted_sum_only fitness assessor will be used.

7.2.34.2.3 roulette_wheel

- Keywords Area
- method
- soga
- replacement_type
- roulette_wheel

Replace population
Specification

Alias: none
Argument(s): none

Description

Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The replacement_type of roulette_wheel or unique_roulette_wheel may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the layer_rank and domination_count, the recommended selector is the below_limit selector. The below_limit replacement will only keep designs that are dominated by fewer than a limiting number of other designs. The replacement_type of favor_feasible is specific to a SOGA. This replacement operator will always prefer a more feasible design to a less feasible one. Beyond that, it favors solutions based on an assigned fitness value which must have been installed by the weighted sum only fitness assessor (see the discussion below).

7.2.34.2.4 unique_roulette_wheel

- Keywords Area
- method
- soga
- replacement_type
- unique_roulette_wheel

Replace population

Specification

Alias: none
Argument(s): none

Description

Replace the population with members selected to continue in the next generation. The pool of potential members is the current population and the current set of offspring. The replacement_type of roulette_wheel or unique_roulette_wheel may be used either with MOGA or SOGA problems however they are not recommended for use with MOGA. Given that the only two fitness assessors for MOGA are the layer_rank and domination_count, the recommended selector is the below_limit selector. The below_limit replacement will only keep designs that are dominated by fewer than a limiting number of other designs. The replacement_type of favor_feasible is specific to a SOGA. This replacement operator will always prefer a more feasible design to a less feasible one. Beyond that, it favors solutions based on an assigned fitness value which must have been installed by the weighted sum only fitness assessor (see the discussion below).

7.2.34.3 convergence_type

- Keywords Area
- method
- soga
• convergence_type

Select the convergence type for JEGA methods

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** average_fitness_tracker

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Convergence Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>best_fitness_tracker</td>
<td>Tracks the best fitness of the population</td>
</tr>
<tr>
<td></td>
<td>average_fitness_tracker</td>
<td></td>
<td>Tracks the average fitness of the population</td>
</tr>
</tbody>
</table>

**Description**

The two JEGA methods use different convergence types, which are described on their respective pages. All the convergence types are modified by the optional keywords percent_change and num_generations.

**7.2.34.3.1 best_fitness_tracker**

• **Keywords Area**
  • method
  • soga
  • convergence_type
  • best_fitness_tracker

Tracks the best fitness of the population

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>percent_change</td>
<td>Define the convergence criterion for JEGA methods</td>
</tr>
</tbody>
</table>
Define the convergence criterion for JEGA methods

**Description**

The **best_fitness_tracker** tracks the best fitness in the population. Convergence occurs after **num_generations** has passed and there has been less than **percent_change** in the best fitness value. The percent change can be as low as 0% in which case there must be no change at all over the number of generations.

**See Also**

These keywords may also be of interest:

- **average_fitness_tracker**

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0.1

**Description**

The **percent_change** is the threshold beneath which convergence is attained whereby it is compared to the metric value computed.

**num_generations**

- **Keywords Area**  
- **method**  
- **soga**  
- **convergence_type**  
- **best_fitness_tracker**  
- **percent_change**

Define the convergence criterion for JEGA methods
7.2 method

Specification

Alias: none

Argument(s): INTEGER

Default: 10

Description

The `num_generations` is the number of generations over which the metric value should be tracked. Convergence will be attained if the recorded metric is below `percent_change` for `num_generations` consecutive generations.

7.2.34.3.2 average_fitness_tracker

- Keywords Area
- method
- soga
- convergence_type
- average_fitness_tracker

Tracks the average fitness of the population

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tr>
<td>Optional</td>
<td>percent_change</td>
<td></td>
<td>Define the convergence criterion for JEGA methods</td>
</tr>
<tr>
<td>Optional</td>
<td>num_generations</td>
<td></td>
<td>Define the convergence criterion for JEGA methods</td>
</tr>
</tbody>
</table>

Description

The `convergence_type` called `average_fitness_tracker` keeps track of the average fitness in a population. If this average fitness does not change more than `percent_change` over some number of generations, `num_generations`, then the solution is reported as converged and the algorithm terminates.

See Also

These keywords may also be of interest:

- best_fitness_tracker
Define the convergence criterion for JEGA methods

**Specification**

**Alias**: none

**Argument(s)**: REAL

**Default**: 0.1

**Description**

The `percent_change` is the threshold beneath which convergence is attained whereby it is compared to the metric value computed.

7.2.34.3.2.2 num_generations

- Keywords Area
- method
- soga
- convergence_type
- average_fitness_tracker
- num_generations

Define the convergence criterion for JEGA methods

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 10

**Description**

The `num_generations` is the number of generations over which the metric value should be tracked. Convergence will be attained if the recorded metric is below `percent_change` for `num_generations` consecutive generations.
7.2.34.4 max_iterations

- Keywords Area
- method
- soga
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.34.5 max_function_evaluations

- Keywords Area
- method
- soga
- max_function_evaluations

Number of function evaluations allowed for optimizers
This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed `max_function_evaluations` evaluations. See also `max_iterations`.

Some optimizers (e.g. `ncsu_direct`) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

**Default Behavior**

Default value is 1000.

---

**Keywords Area**

**method**

**soga**

**scaling**

Turn on scaling for variables, responses, and constraints

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords \texttt{scale\_types} and/or \texttt{*scales} in the \texttt{variables} and \texttt{responses} blocks. When the \texttt{scaling} keyword is omitted from \texttt{method}, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a \texttt{*scales} keyword, which gives characteristic values (divisors)
- a \texttt{*scale\_types} keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in \texttt{responses} for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

\textbf{Scale Types}

There are three scale types:

1. \texttt{value} - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. \texttt{auto} - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component
   
   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. \texttt{log} - logarithmic scaling
   First, any characteristic values from the optional \texttt{*_scales} specification are applied. Then logarithm base 10 scaling is applied.
   
   Logarithmic scaling is not available for linear constraints.
   
   When continuous design variables are log scaled, linear constraints are not allowed.

\textbf{Scales}

The \texttt{*scales} keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding \texttt{*scale\_types}, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- \texttt{auto, log} - optional
- \texttt{value} - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL\_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
 objective_functions 3
 sense "maximize"
 primary_scale_types = "value" "value" "value"
 primary_scales = 1 1 100

responses
 objective_functions 3
 sense "maximize"
 primary_scale_types = "value" "value" "value"
 primary_scales = 1 1 100
```

7.2.34.7 population_size

- Keywords Area
- method
- soga
- population_size

Set the initial population size in JEGA methods

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 50

**Description**

The number of designs in the initial population is specified by the `population_size`. Note that the `population_size` only sets the size of the initial population. The population size may vary in the JEGA methods according to the type of operators chosen for a particular optimization run.

Also note that some initializers may not strictly respect the initial population size. See the SOGA `flat_file` or MOGA `flat_file` initializer for an example.

7.2.34.8 log_file

- Keywords Area
- method
- soga
- log_file

Specify the name of a log file
7.2 method

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** JEGAGlobal.log

**Description**

New as of JEGA v2.0 is the introduction of the log_file specification. JEGA now uses a logging library to output messages and status to the user. JEGA can be configured at build time to log to both the console window and a text file, one or the other, or neither. The log_file input is a string name of a file into which to log. If the build was configured without file logging in JEGA, this input is ignored. If file logging is enabled and no log_file is specified, the default file name of JEGAGlobal.log is used.

7.2.34.9 print_each_pop

- **Keywords Area**
- **method**
- **soga**
- **print_each_pop**

Print every population to a population file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** No printing

**Description**

New to JEGA v2.0 is the introduction of the print_each_pop specification. It serves as a flag and if supplied, the population at each generation will be printed to a file named "population<GEN#>.dat" where <GEN#> is the number of the current generation.

7.2.34.10 initialization_type

- **Keywords Area**
- **method**
- **soga**
- **initialization_type**

Specify how to initialize the population
Specification

**Alias:** none

**Argument(s):** none

**Default:** unique_random

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group Initialization Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td><strong>Required</strong> (Choose One)</td>
<td></td>
<td>simple_random</td>
<td>Create random initial solutions</td>
</tr>
<tr>
<td>unique_random</td>
<td>Create random initial solutions, but enforce uniqueness (default)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>flat_file</td>
<td>Read initial solutions from file</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The `initialization_type` defines how the initial population is created for the GA. There are three types:

1. `simple_random`
2. `unique_random` (default)
3. `flat_file`

Setting the size for the `flat_file` initializer has the effect of requiring a minimum number of designs to create. If this minimum number has not been created once the files are all read, the rest are created using the `unique_random` initializer and then the `simple_random` initializer if necessary.

7.2.34.10.1 `simple_random`

- Keywords Area
- method
- soga
- initialization_type
- simple_random

Create random initial solutions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**
simple_random creates initial solutions with random variable values according to a uniform random number distribution. It gives no consideration to any previously generated designs.
7.2.34.10.2 unique_random

- Keywords Area
- method
- soga
- initialization_type
- unique_random

Create random initial solutions, but enforce uniqueness (default)

Specification

Alias: none
Argument(s): none

Description

unique_random is the same as simple_random, except that when a new solution is generated, it is checked against the rest of the solutions. If it duplicates any of them, it is rejected.

7.2.34.10.3 flat_file

- Keywords Area
- method
- soga
- initialization_type
- flat_file

Read initial solutions from file

Specification

Alias: none
Argument(s): STRING

Description

flat_file allows the initial population to be read from a flat file. If flat_file is specified, a file name must be given.

Variables can be delimited in the flat file in any way you see fit with a few exceptions. The delimiter must be the same on any given line of input with the exception of leading and trailing whitespace. So a line could look like: 1.1, 2.2, 3.3 for example but could not look like: 1.1, 2.2 3.3. The delimiter can vary from line to line within the file which can be useful if data from multiple sources is pasted into the same input file. The delimiter can be any string that does not contain any of the characters .+-dDeE or any of the digits 0-9. The input will be read until the end of the file. The algorithm will discard any configurations for which it was unable to retrieve at least the number of design variables. The objective and constraint entries are not required but if ALL are present, they will be recorded and the design will be tagged as evaluated so that evaluators may choose not to re-evaluate them.
Setting the size for this initializer has the effect of requiring a minimum number of designs to create. If this minimum number has not been created once the files are all read, the rest are created using the unique_random initializer and then the simple_random initializer if necessary. If more designs are found in the initialization files than specified by the initial size, then the initial size is ignored and all designs read out of the files are kept in the initial population.

### 7.2.34.11 crossover_type

- **Keywords Area**
  - method
  - soga
  - crossover_type

Select a crossover type for JEGA methods

**Specification**

- **Alias:** none
- **Argument(s):** none
- **Default:** shuffle_random

**Child Keywords:**

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<th>Required/Optional</th>
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<th>Dakota Keyword Description</th>
</tr>
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<td>Crossover Type (Group 1)</td>
<td>multi_point_binary</td>
<td>Use bit switching for crossover events</td>
</tr>
<tr>
<td></td>
<td></td>
<td>multi_point_-parameterized_binary</td>
<td>Use bit switching to crossover each design variable</td>
</tr>
<tr>
<td></td>
<td></td>
<td>multi_point_real</td>
<td>Perform crossover in real valued genome</td>
</tr>
<tr>
<td></td>
<td></td>
<td>shuffle_random</td>
<td>Perform crossover by choosing design variable(s)</td>
</tr>
</tbody>
</table>

**Optional**

|                  | crossover_rate | Specify the probability of a crossover event |

**Description**

There are many crossover types available. **multi_point_binary** crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). **multi_point_parameterized_binary** crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. **multi_point_real** crossover performs a variable switching crossover routing at N crossover points in the real real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by $10^{36}$ and then truncating. Note that this
assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

The final crossover type is shuffle_random. This crossover type performs crossover by choosing design variables at random from a specified number of parents enough times that the requested number of children are produced. For example, consider the case of 3 parents producing 2 children. This operator would go through and for each design variable, select one of the parents as the donor for the child. So it creates a random shuffle of the parent design variable values. The relative numbers of children and parents are controllable to allow for as much mixing as desired. The more parents involved, the less likely that the children will wind up exact duplicates of the parents.

All crossover types take a **crossover_rate**. The crossover rate is used to calculate the number of crossover operations that take place. The number of crossovers is equal to the rate * population_size.

### 7.2.34.11.1 mult\_point\_binary

- **Keywords Area**
- **method**
- **soga**
- **crossover\_type**
- **mult\_point\_binary**

Use bit switching for crossover events

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

There are many crossover types available. **mult\_point\_binary** crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). **mult\_point\_parameterized\_binary** crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. **mult\_point\_real** crossover performs a variable switching crossover routine at N crossover points in the real real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by 10^6 and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

### 7.2.34.11.2 mult\_point\_parameterized\_binary

- **Keywords Area**
- **method**
- **soga**
- **crossover\_type**
- **mult\_point\_parameterized\_binary**

Use bit switching to crossover each design variable
There are many crossover types available. `multi_point_binary` crossover requires an integer number, N, of crossover points. This crossover type performs a bit switching crossover at N crossover points in the binary encoded genome of two designs. Thus, crossover may occur at any point along a solution chromosome (in the middle of a gene representing a design variable, for example). `multi_point_parameterized_binary` crossover is similar in that it performs a bit switching crossover routine at N crossover points. However, this crossover type performs crossover on each design variable individually. So the individual chromosomes are crossed at N locations. `multi_point_real` crossover performs a variable switching crossover routing at N crossover points in the real valued genome of two designs. In this scheme, crossover only occurs between design variables (chromosomes). Note that the standard solution chromosome representation in the JEGA algorithm is real encoded and can handle integer or real design variables. For any crossover types that use a binary representation, real variables are converted to long integers by multiplying the real number by $10^6$ and then truncating. Note that this assumes a precision of only six decimal places. Discrete variables are represented as integers (indices within a list of possible values) within the algorithm and thus require no special treatment by the binary operators.

Perform crossover in real valued genome
Perform crossover by choosing design variable(s)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>num_parents</td>
<td>num_parents</td>
<td>Number of parents in random shuffle crossover</td>
</tr>
<tr>
<td>Optional</td>
<td>num_offspring</td>
<td>num_offspring</td>
<td>Number of offspring in random shuffle crossover</td>
</tr>
</tbody>
</table>

**Description**

The final crossover type is **shuffle_random**. This crossover type performs crossover by choosing design variables at random from a specified number of parents enough times that the requested number of children are produced. For example, consider the case of 3 parents producing 2 children. This operator would go through and for each design variable, select one of the parents as the donor for the child. So it creates a random shuffle of the parent design variable values. The relative numbers of children and parents are controllable to allow for as much mixing as desired. The more parents involved, the less likely that the children will wind up exact duplicates of the parents.

7.2.34.11.4.1 num_parents

**Alias:** none

**Argument(s):** none

**Child Keywords:**

Number of parents in random shuffle crossover
Keywords Area

Specification

Alias: none
Argument(s): INTEGER
Default: 2

Description

Number of parents in random shuffle crossover

7.2.34.11.4.2 num_offspring

• Keywords Area
• method
• soga
• crossover_type
• shuffle_random
• num_offspring

Number of offspring in random shuffle crossover

Specification

Alias: none
Argument(s): INTEGER
Default: 2

Description

Number of offspring in random shuffle crossover

7.2.34.11.5 crossover_rate

• Keywords Area
• method
• soga
• crossover_type
• crossover_rate

Specify the probability of a crossover event

Specification

Alias: none
Argument(s): REAL
Default: 0.8
The `crossover_type` controls what approach is employed for combining parent genetic information to create offspring, and the `crossover_rate` specifies the probability of a crossover operation being performed to generate a new offspring. The SCOLIB EA method supports three forms of crossover, `two_point`, `blend`, and `uniform`, which generate a new individual through combinations of two parent individuals. Two-point crossover divides each parent into three regions, where offspring are created from the combination of the middle region from one parent and the end regions from the other parent. Since the SCOLIB EA does not utilize bit representations of variable values, the crossover points only occur on coordinate boundaries, never within the bits of a particular coordinate. Uniform crossover creates offspring through random combination of coordinates from the two parents. Blend crossover generates a new individual randomly along the multidimensional vector connecting the two parents.

### 7.2.34.12 mutation_type

- **Keywords Area**
  - method
  - soga
  - mutation_type

Select a mutation type for JEGA methods

#### Specification

**Alias:** none

**Argument(s):** none

**Default:** `replace_uniform`

#### Child Keywords:

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<th>Dakota Keyword</th>
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<td>replace_uniform</td>
<td>Mutate by flipping a random bit</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_normal</td>
<td>Set mutation offset to use a normal distribution</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_cauchy</td>
<td>Use a Cauchy distribution for the mutation offset</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_uniform</td>
<td>Set mutation offset to use a uniform distribution</td>
</tr>
<tr>
<td>Optional</td>
<td>mutation_rate</td>
<td></td>
<td>Set probability of a mutation</td>
</tr>
</tbody>
</table>

#### Description

Five mutation types are available for selection by keyword: `replace_uniform`, `bit_random`, `offset_cauchy`, `offset_normal`, and `offset_uniform`. They are described in greater detail on their respective keyword pages.
The offset_\* mutators all act by adding a random "offset" to a variable value. The random amount has a mean of zero in all cases. The size of the offset is controlled using the \texttt{mutation\_scale} keyword, which is interpreted differently for each offset_\* type.

The rate of mutations for all types is controlled using the \texttt{mutation\_rate}. The rate is applied differently in each mutation_type.

7.2.34.12.1 bit_random

- Keywords Area
- method
- soga
- mutation_type
- bit_random

Mutate by flipping a random bit

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The bit_random mutator introduces random variation by first converting a randomly chosen variable of a randomly chosen design into a binary string. It then flips a randomly chosen bit in the string from a 1 to a 0 or visa versa. In this mutation scheme, the resulting value has more probability of being similar to the original value.

7.2.34.12.2 replace_uniform

- Keywords Area
- method
- soga
- mutation_type
- replace_uniform

Use uniformly distributed value over range of parameter

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

replace_uniform introduces random variation by first randomly choosing a design variable of a randomly selected design and reassigning it to a random valid value for that variable. No consideration of the current value is given when determining the new value.
Set mutation offset to use a normal distribution

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_scale</td>
<td>Scales mutation across range of parameter</td>
</tr>
</tbody>
</table>

Description

The offset_normal mutator introduces random variation by adding a Gaussian random amount to a variable value. The random amount has a standard deviation dependent on the mutation_scale.

Scales mutation across range of parameter

Specification

Alias: none

Argument(s): REAL

Default: 0.15
The mutation_scale is a fraction in the range [0, 1] and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected mutation_type. For offset_NORMAL and offset_cauchy, mutation_scale is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For offset_uniform, the range of possible deviation amounts is +/- 1/2 * (mutation_scale * variable range).

7.2.34.12.4 offset_cauchy

- Keywords Area
- method
- soga
- mutation_type
- offset_cauchy

Use a Cauchy distribution for the mutation offset

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_scale</td>
<td>Scales mutation across range of parameter</td>
</tr>
</tbody>
</table>

**Description**

The offset_cauchy mutator introduces random variation by adding a Cauchy random amount to a variable value. The random amount has a standard deviation dependent on the mutation_scale.

7.2.34.12.4.1 mutation_scale

- Keywords Area
- method
- soga
- mutation_type
- offset_cauchy
- mutation_scale

Scales mutation across range of parameter

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0.15
Description

The mutation_scale is a fraction in the range [0, 1] and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected mutation_type. For offset-normal and offset_cauchy, mutation_scale is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For offset_uniform, the range of possible deviation amounts is +/- 1/2 * (mutation_scale * variable range).

7.2.34.12.5 offset_uniform

- Keywords Area
- method
- soga
- mutation_type
- offset_uniform

Set mutation offset to use a uniform distribution

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_scale</td>
<td>Scales mutation across range of parameter</td>
</tr>
</tbody>
</table>

Description

The offset_uniform mutator introduces random variation by adding a uniform random amount to a variable value. The random amount depends on the mutation_scale.

7.2.34.12.5.1 mutation_scale

- Keywords Area
- method
- soga
- mutation_type
- offset_uniform
- mutation_scale

Scales mutation across range of parameter

Specification

Alias: none

Argument(s): REAL

Default: 0.15
The mutation_scale is a fraction in the range \([0, 1]\) and is meant to help control the amount of variation that takes place when a variable is mutated. Its behavior depends on the selected mutation_type. For offset_normal and offset_cauchy, mutation_scale is multiplied by the range of the variable being mutated to obtain the standard deviation of the offset. For offset_uniform, the range of possible deviation amounts is \(+/-(mutation_scale \times variable\ range)\).

7.2.34.12.6 mutation_rate

- Keywords Area
- method
- soga
- mutation_type
- mutation_rate

Set probability of a mutation

Specification

Alias: none

Argument(s): REAL

Default: 0.08

Description

All mutation types have a mutation_rate, which controls the number of mutations performed. For replace_uniform and all the offset_* types, the number of mutations performed is the product of mutation_rate and population_size. For bit_random, it's the product of the mutation_rate, number of design variables, and population size.

7.2.34.13 seed

- Keywords Area
- method
- soga
- seed

Seed of the random number generator

Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)
Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.34.14 convergence_tolerance

- Keywords Area
- method
- soga
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.
Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.34.15 model_pointer

**Keywords Area**

- **method**
- **soga**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

*Alias:* none

*Argument(s):* STRING

*Default:* method use of last model parsed (or use of default model if none parsed)

**Description**

The **model_pointer** is used to specify which **model** block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a **model** block in the Dakota input file that has a corresponding **id_model** with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a **model_pointer** for each method is imperative.

See **block_pointer** for details about pointers.
Examples

```dakota
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
    sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians
```

7.2.35 coliny_pattern_search

- Keywords Area
- method
- coliny_pattern_search

Pattern search, derivative free optimization method

Topics

This keyword is related to the topics:

- package_scolib
- package_coliny
- global_optimization_methods

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td></td>
<td>Use a simple weighted penalty to manage feasibility</td>
</tr>
<tr>
<td>Optional</td>
<td>no_expansion</td>
<td></td>
<td>Don't allow expansion of the search pattern</td>
</tr>
<tr>
<td>Optional</td>
<td>expand_after_-success</td>
<td></td>
<td>Set the factor by which a search pattern can be expanded</td>
</tr>
<tr>
<td>Optional</td>
<td>pattern_basis</td>
<td></td>
<td>Pattern basis selection</td>
</tr>
<tr>
<td>Optional</td>
<td>stochastic</td>
<td></td>
<td>Generate trial points in random order</td>
</tr>
<tr>
<td>Optional</td>
<td>total_pattern_size</td>
<td></td>
<td>Total number of points in search pattern</td>
</tr>
<tr>
<td>Optional</td>
<td>exploratory_moves</td>
<td></td>
<td>Exploratory moves selection</td>
</tr>
<tr>
<td>Optional</td>
<td>synchronization</td>
<td></td>
<td>Select how Dakota schedules a batch of concurrent function evaluations in a parallel algorithm</td>
</tr>
<tr>
<td>Optional</td>
<td>contraction_factor</td>
<td></td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td>constraint_penalty</td>
<td></td>
<td>Multiplier for the penalty function</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_delta</td>
<td></td>
<td>Initial step size for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>variable_tolerance</td>
<td></td>
<td>Step length-based stopping criteria for derivative-free optimizers</td>
</tr>
</tbody>
</table>
Pattern search techniques are nongradient-based optimization methods which use a set of offsets from the current iterate to locate improved points in the design space.

See the page package_scolib for important information regarding all SCOLIB methods

colin_pattern_search supports concurrency up to the size of the search pattern

Traditional pattern search methods search with a fixed pattern of search directions to try to find improvements to the current iterate. The SCOLIB pattern search methods generalize this simple algorithmic strategy to enable control of how the search pattern is adapted, as well as how each search pattern is evaluated. The stochastic and synchronization specifications denote how the trial points are evaluated. The stochastic specification indicates that the trial points are considered in a random order. For parallel pattern search, synchronization dictates whether the evaluations are scheduled using a blocking scheduler or a nonblocking scheduler. In the blocking case, all points in the pattern are evaluated (in parallel), and if the best of these trial points is an improving point, then it becomes the next iterate. These runs are reproducible, assuming use of the same seed in the stochastic case. In the nonblocking case, all points in the pattern may not be evaluated, since the first improving point found becomes the next iterate. Since the algorithm steps will be subject to parallel timing variabili-
ties, these runs will not generally be repeatable. The synchronization specification has similar connotations for sequential pattern search. If blocking is specified, then each sequential iteration terminates after all trial points have been considered, and if nonblocking is specified, then each sequential iteration terminates after the first improving trial point is evaluated. In this release, both blocking and nonblocking specifications result in blocking behavior (except in the case where exploratory_moves below is set to adaptive_pattern). Nonblocking behavior will be re-enabled after some underlying technical issues have been resolved.

The particular form of the search pattern is controlled by the pattern_basis specification. If pattern_basis is coordinate basis, then the pattern search uses a plus and minus offset in each coordinate direction, for a total of \(2n\) function evaluations in the pattern. This case is depicted in Figure 5.3 for three coordinate dimensions.

![Figure 7.1: Depiction of coordinate pattern search algorithm](image_url)

If pattern_basis is simplex, then pattern search uses a minimal positive basis simplex for the parameter space, for a total of \(n+1\) function evaluations in the pattern. Note that the simplex pattern basis can be used for unbounded problems only. The total_pattern_size specification can be used to augment the basic coordinate and simplex patterns with additional function evaluations, and is particularly useful for parallel load balancing. For example, if some function evaluations in the pattern are dropped due to duplication or bound constraint interaction, then the total_pattern_size specification instructs the algorithm to generate new offsets to bring the total number of evaluations up to this consistent total.

The exploratory_moves specification controls how the search pattern is adapted. (The search pattern can be adapted after an improving trial point is found, or after all trial points in a search pattern have been found to be unimproving points.) The following exploratory moves selections are supported by SCOLIB:

- The basic_pattern case is the simple pattern search approach, which uses the same pattern in each iteration.

- The multi_step case examines each trial step in the pattern in turn. If a successful step is found, the pattern search continues examining trial steps about this new point. In this manner, the effects of multiple successful steps are cumulative within a single iteration. This option does not support any parallelism and will result in a serial pattern search.
The **adaptive_pattern** case invokes a pattern search technique that adaptively rescales the different search directions to maximize the number of redundant function evaluations. See [46] for details of this method. In preliminary experiments, this method had more robust performance than the standard **basic_pattern** case in serial tests. This option supports a limited degree of parallelism. After successful iterations (where the step length is not contracted), a parallel search will be performed. After unsuccessful iterations (where the step length is contracted), only a single evaluation is performed.

The **initial_delta** and **variable_tolerance** specifications provide the initial offset size and the threshold size at which to terminate the algorithm. For any dimension that has both upper and lower bounds, this step length will be internally rescaled to provide search steps of length \( \text{initial_delta} \times \text{range} \times 0.1 \). This rescaling does not occur for other dimensions, so search steps in those directions have length \( \text{initial_delta} \). Note that the factor of 0.1 in the rescaling could result in an undesirably small initial step. This can be offset by providing a large **initial_delta**.

In general, pattern search methods can expand and contract their step lengths. SCOLIB pattern search methods contract the step length by the value **contraction_factor**, and they expand the step length by the value \((1/\text{contraction_factor})\). The **expand_after_success** control specifies how many successful objective function improvements must occur with a specific step length prior to expansion of the step length, whereas the **no_expansion** flag instructs the algorithm to forgo pattern expansion altogether.

Finally, constraint infeasibility can be managed in a somewhat more sophisticated manner than the simple weighted penalty function. If the **constant_penalty** specification is used, then the simple weighted penalty scheme described above is used. Otherwise, the constraint penalty is adapted to the value \( \text{constraint_penalty}/L \), where \( L \) is the the smallest step length used so far.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the **hdf5** keyword, this method writes the following results to HDF5:

- **Best Parameters**
- **Best Objective Functions** (when **objective_functions** are specified)
- **Best Nonlinear Constraints**
- **Calibration** (when **calibration_terms** are specified)

**See Also**

These keywords may also be of interest:

- **coliny_beta**
- **coliny_direct**
- **coliny_cobyla**
- **coliny_ea**
- **coliny_solis_wets**
Specification

**Alias:** none

**Argument(s):** none

**Default:** algorithm dynamically adapts the constraint penalty

Description

Finally, constraint infeasibility can be managed in a somewhat more sophisticated manner than the simple weighted penalty function. If the `constant_penalty` specification is used, then the simple weighted penalty scheme described above is used. Otherwise, the constraint penalty is adapted to the value `constraint_penalty/L`, where L is the the smallest step length used so far.

7.2.35.2 no_expansion

- **Keywords Area**
- **method**
- **coliny_pattern_search**
- **no_expansion**

Don’t allow expansion of the search pattern

Specification

**Alias:** none

**Argument(s):** none

**Default:** algorithm may expand pattern size

Description

In general, pattern search methods can expand and contract their step lengths. SCOLIB pattern search methods contract the step length by the value `contraction_factor`, and they expand the step length by the value `(1/contraction_factor)`. The `expand_after_success` control specifies how many successful objective function improvements must occur with a specific step length prior to expansion of the step length, whereas the `no_expansion` flag instructs the algorithm to forgo pattern expansion altogether.

7.2.35.3 expand_after_success

- **Keywords Area**
- **method**
- **coliny_pattern_search**
- **expand_after_success**

Set the factor by which a search pattern can be expanded
Specification

Alias: none
Argument(s): INTEGER
Default: 5

Description

In general, pattern search methods can expand and contract their step lengths. SCOLIB pattern search methods contract the step length by the value \( \text{contraction} \_\text{factor} \), and they expand the step length by the value \((1/\text{contraction} \_\text{factor})\). The \text{expand} \_\text{after} \text{success} control specifies how many successful objective function improvements must occur with a specific step length prior to expansion of the step length, whereas the \text{no} \_\text{expansion} flag instructs the algorithm to forgo pattern expansion altogether.

7.2.35.4 pattern_basis

- Keywords Area
- method
- coliny_pattern_search
- pattern_basis

Pattern basis selection

Specification

Alias: none
Argument(s): none
Default: coordinate

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Pattern Basis Type (Group 1)</td>
<td>coordinate</td>
<td>Use coordinate directions as search pattern</td>
</tr>
<tr>
<td></td>
<td>simplex</td>
<td></td>
<td>Use a minimal simplex for the search pattern</td>
</tr>
</tbody>
</table>

Description

The particular form of the search pattern is controlled by the \text{pattern} \_\text{basis} specification. If \text{pattern} \_\text{basis} is \text{coordinate} basis, then the pattern search uses a plus and minus offset in each coordinate direction, for a total of \( 2n \) function evaluations in the pattern. This case is depicted in Figure 5.3 for three coordinate dimensions.

7.2.35.4.1 coordinate

- Keywords Area
- method
- coliny_pattern_search
Use coordinate directions as search pattern

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The particular form of the search pattern is controlled by the `pattern_basis` specification. If `pattern_basis` is `coordinate` basis, then the pattern search uses a plus and minus offset in each coordinate direction, for a total of $2n$ function evaluations in the pattern. This case is depicted in Figure 5.3 for three coordinate dimensions.

**7.2.35.4.2 simplex**

- `Keywords Area`
- `method`
- `coliny_pattern_search`
- `pattern_basis`
- `simplex`

Use a minimal simplex for the search pattern

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

If `pattern_basis` is `simplex`, then pattern search uses a minimal positive basis simplex for the parameter space, for a total of $n+1$ function evaluations in the pattern. Note that the `simplex` pattern basis can be used for unbounded problems only. The `total_pattern_size` specification can be used to augment the basic coordinate and simplex patterns with additional function evaluations, and is particularly useful for parallel load balancing. For example, if some function evaluations in the pattern are dropped due to duplication or bound constraint interaction, then the `total_pattern_size` specification instructs the algorithm to generate new offsets to bring the total number of evaluations up to this consistent total.

**7.2.35.5 stochastic**

- `Keywords Area`
- `method`
- `coliny_pattern_search`
- `stochastic`

Generate trial points in random order
7.2 method

Specification

Alias: none
Argument(s): none

Description

Traditional pattern search methods search with a fixed pattern of search directions to try to find improvements to the current iterate. The SCOLIB pattern search methods generalize this simple algorithmic strategy to enable control of how the search pattern is adapted, as well as how each search pattern is evaluated. The stochastic and synchronization specifications denote how the trial points are evaluated. The stochastic specification indicates that the trial points are considered in a random order. For parallel pattern search, synchronization dictates whether the evaluations are scheduled using a blocking scheduler or a nonblocking scheduler (i.e.,

7.2.35.6 total_pattern_size

- Keywords Area
- method
- coliny_pattern_search
- total_pattern_size

Total number of points in search pattern

Specification

Alias: none
Argument(s): INTEGER
Default: no augmentation of basic pattern

Description

If pattern_basis is simplex, then pattern search uses a minimal positive basis simplex for the parameter space, for a total of n+1 function evaluations in the pattern. Note that the simplex pattern basis can be used for unbounded problems only. The total_pattern_size specification can be used to augment the basic coordinate and simplex patterns with additional function evaluations, and is particularly useful for parallel load balancing. For example, if some function evaluations in the pattern are dropped due to duplication or bound constraint interaction, then the total_pattern_size specification instructs the algorithm to generate new offsets to bring the total number of evaluations up to this consistent total.

7.2.35.7 exploratory_moves

- Keywords Area
- method
- coliny_pattern_search
- exploratory_moves

Exploratory moves selection
### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** basic_pattern  

#### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Exploration Mode (Group 1)</td>
<td>multi_step</td>
<td>Examine trial step around successful new point</td>
</tr>
<tr>
<td>adaptive_pattern</td>
<td>Adaptively rescale search directions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>basic_pattern</td>
<td>Use the same search pattern every iteration</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

#### Description

The exploratory_moves specification controls how the search pattern is adapted. (The search pattern can be adapted after an improving trial point is found, or after all trial points in a search pattern have been found to be unimproving points.) The following exploratory moves selections are supported by SCOLIB:

**7.2.35.7.1 multi_step**

- Keywords Area
- method
- coliny_pattern_search
- exploratory_moves
- multi_step

Examine trial step around successful new point

#### Specification

**Alias:** none  
**Argument(s):** none  

#### Description

The **multi_step** case examines each trial step in the pattern in turn. If a successful step is found, the pattern search continues examining trial steps about this new point. In this manner, the effects of multiple successful steps are cumulative within a single iteration. This option does not support any parallelism and will result in a serial pattern.
Adaptively rescale search directions

Specification

Alias: none

Argument(s): none

Description

The adaptive_pattern case invokes a pattern search technique that adaptively rescales the different search directions to maximize the number of redundant function evaluations. See [46] for details of this method. In preliminary experiments, this method had more robust performance than the standard basic_pattern case in serial tests. This option supports a limited degree of parallelism. After successful iterations (where the step length is not contracted), a parallel search will be performed. After unsuccessful iterations (where the step length is contracted), only a single evaluation is performed.

7.2.35.7.3 basic_pattern

Use the same search pattern every iteration

Specification

Alias: none

Argument(s): none

Description

The basic_pattern case is the simple pattern search approach, which uses the same pattern in each iteration.

7.2.35.8 synchronization

Select how Dakota schedules a batch of concurrent function evaluations in a parallel algorithm
Specification

**Alias:** none

**Argument(s):** none

**Default:** method specific

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required</strong> <em>(Choose One)</em></td>
<td></td>
<td>blocking</td>
<td>Complete concurrent evaluation of all evaluations within a batch</td>
</tr>
<tr>
<td></td>
<td></td>
<td>nonblocking</td>
<td>A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm</td>
</tr>
</tbody>
</table>

**Description**

The synchronization specification can be used to specify the use of either blocking or nonblocking schedulers, where the former must block until all evaluations are completed and the latter does not block, returning only the partial set that has been completed for a particular call to the scheduler.

7.2.35.8.1 blocking

- [Keywords Area](#)
- [method](#)
- [coliny_pattern_search](#)
- [synchronization](#)
- [blocking](#)

Complete concurrent evaluation of all evaluations within a batch

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

In the **blocking** case, all of the concurrent evaluations in the batch must be completed and all results are returned together to the algorithm.

These runs are reproducible, assuming use of the same seed in cases with random number generation.
A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

In the nonblocking case, the set of running evaluations is queried for completion and a partial set of results is returned to an asynchronous algorithm, which can then generate new evaluations based on these partial results.

In all cases, the algorithm steps will be subject to parallel timing variabilities / race conditions such that these runs will not generally be repeatable regardless of seed specification.

**7.2.35.9 contraction_factor**

Amount by which step length is rescaled

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0.5

**Description**

For pattern search methods, `contraction_factor` specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is $1/\text{contraction\_factor}$.
7.2.35.10 constraint_penalty

- Keywords Area
- method
- coliny_pattern_search
- constraint_penalty

Multiplier for the penalty function

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0

**Description**

Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds \( \text{constraint\_penalty} \times \) the sum of squares of the constraint violations to the objective function. The default value of \( \text{constraint\_penalty} \) is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.

7.2.35.11 initial_delta

- Keywords Area
- method
- coliny_pattern_search
- initial_delta

Initial step size for derivative-free optimizers

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0 (COBYLA), 0.1*range (PS, SW)

**Description**

The \( \text{initial\_delta} \) keyword defines the size of the first search step in derivative-free optimization methods, specifically \( \text{asynch\_pattern\_search}, \text{coliny\_cobyla}, \text{coliny\_pattern\_search}, \text{coliny\_solis\_wets}, \text{and mesh\_adaptive\_search} \). It is applied in an absolute sense to all search directions.

**Default Behavior**

The default value is 1.0.

**Usage Tips**
It is recommended that initial_delta be the approximate distance from the initial point to the solution. If this distance is unknown, it is advisable to err on the side of choosing an initial_delta that is too large or to not specify it. Relative application of initial_delta is not available unless the user scales the problem accordingly.

Examples

Three example method input blocks appear below.

For asynch_pattern_search:

```plaintext
method
  asynch_pattern_search
  initial_delta = .5
  contraction_factor = 0.25
  merit_function merit1_smooth
  smoothing_factor = 1.0
  constraint_tolerance = 1.e-6
```

For coliny_pattern_search:

```plaintext
method
  coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
```

For mesh_adaptive_search

```plaintext
method
  mesh_adaptive_search
  initial_delta = 2.0
  seed = 1234
```

7.2.35.12 variable_tolerance

- Keywords Area
- method
  - coliny_pattern_search
  - variable_tolerance

Step length-based stopping criteria for derivative-free optimizers

Specification

Alias: none

Argument(s): REAL

Default: 1.0e-4 (COBYLA), 1.0e-5 (PS), 1.0e-6 (SW)

Description

The variable_tolerance keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to asynch_pattern_search, coliny_cobyla, coliny_pattern_search, coliny_solis_wets, and mesh_adaptive_search.
Default Behavior

The default value varies according to method as follows:

- `asynch_pattern_search`: 1.0e-2
- `coliny_cobyla`: 1.0e-4
- `coliny_pattern_search`: 1.0e-5
- `coliny_solis_wets`: 1.0e-6
- `mesh_adaptive_search`: 1.0e-6

Usage Tips

It is recommended that `variable_tolerance` be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.

Examples

Three example method input blocks appear below.

For `asynch_pattern_search`:

```plaintext
method
  asynch_pattern_search
  contraction_factor = 0.25
  variable_tolerance = 1.e-4
  solution_target = 1.e-6
  max_function_evaluations 500
  constraint_tolerance 1.e-6
```

For `coliny_pattern_search`:

```plaintext
method
  coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
```

For `mesh_adaptive_search`

```plaintext
method
  mesh_adaptive_search
  variable_tolerance = 0.01
  seed = 1234
```

Stopping criteria based on objective function value
7.2 method

Specification

Alias: solution_accuracy

Argument(s): REAL

Default: -DBL_MAX

Description

solution_target is a termination criterion. The algorithm will terminate when the function value falls below solution_target.

7.2.35.14 seed

- Keywords Area
- method
- coliny_pattern_search
- seed

Seed of the random number generator

Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```
7.2.35.15  show_misc_options

    • Keywords Area
    • method
    • coliny_pattern_search
    • show_misc_options

Show algorithm parameters not exposed in Dakota input

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no dump of specification options

**Description**

All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputing additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification: they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

7.2.35.16  misc_options

    • Keywords Area
    • method
    • coliny_pattern_search
    • misc_options

Set method options not available through Dakota spec

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** no misc options

**Description**

All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputing additional settings supported by the SCOLIB
methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

7.2.35.17 max_iterations

- Keywords Area
- method
- coliny_pattern_search
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max_function_evaluations

7.2.35.18 convergence_tolerance

- Keywords Area
- method
- coliny_pattern_search
• convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

• DOT: relative tolerance that must be satisfied for two consecutive iterations
• NL2SOL: See nl2sol
• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.35.19 max_function_evaluations

• Keywords Area

• method

• coliny_pattern_search

• max_function_evaluations

Number of function evaluations allowed for optimizers
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.35.20 scaling

- Keywords Area
- method
- coliny_pattern_search
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `sc*ales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `sc*ales` keyword, which gives characteristic values (divisors)
- a `sc*ale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

### Scale Types

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `sc*ales` specification are applied. Then logarithm base 10 scaling is applied.
   
   Logarithmic scaling is not available for linear constraints.
   
   When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The `sc*ales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `sc*ale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- `auto, log` - optional
- `value` - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than `1.0e10*DBL_MIN`. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.35.21  model_pointer

- Keywords Area
- method
- coliny_pattern_search
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.
Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
 polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
 lower_bounds = 0. 0.
 upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

7.2.36 coliny_solis_wets

- Keywords Area
- method
- coliny_solis_wets

Simple greedy local search method

Topics

This keyword is related to the topics:

- package_scolib
- package_coliny
### Specification

**Alias:** none  
**Argument(s):** none

#### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>contract_after_-failure</td>
<td>no_expansion</td>
<td>The number of unsuccessful cycles prior to contraction.</td>
</tr>
<tr>
<td>Optional</td>
<td>no_expansion</td>
<td>expand_after_-success</td>
<td>Don't allow expansion of the search pattern</td>
</tr>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td>contraction_factor</td>
<td>Set the factor by which a search pattern can be expanded</td>
</tr>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td>initial_delta</td>
<td>Use a simple weighted penalty to manage feasibility</td>
</tr>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td>variable_tolerance</td>
<td>Amount by which step length is rescaled</td>
</tr>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td>solution_target</td>
<td>Multiplier for the penalty function</td>
</tr>
<tr>
<td>Optional</td>
<td>constant_penalty</td>
<td>seed</td>
<td>Initial step size for derivative-free optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>show_misc_options</td>
<td>show_misc_options</td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>Optional</td>
<td>show_misc_options</td>
<td>misc_options</td>
<td>Set method options not available through Dakota spec</td>
</tr>
</tbody>
</table>

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Optional  
max_iterations  
Number of iterations allowed for optimizers and adaptive UQ methods

Optional  
convergence_-tolerance  
Stopping criterion based on objective function or statistics convergence

Optional  
max_function_-evaluations  
Number of function evaluations allowed for optimizers

Optional  
scaling  
Turn on scaling for variables, responses, and constraints

Optional  
model_pointer  
Identifier for model block to be used by a method

Description

The Solis-Wets method is a simple greedy local search heuristic for continuous parameter spaces. Solis-Wets generates trial points using a multivariate normal distribution, and unsuccessful trial points are reflected about the current point to find a descent direction.

See the page package_scolib for important information regarding all SCOLIB methods

colini_solis_wets is inherently serial, no concurrency is used.

These specifications have the same meaning as corresponding specifications for coliny_pattern_search. Please see that page for specification details.

In particular, coliny_solis_wets supports dynamic rescaling of the step length, and dynamic rescaling of the constraint penalty. The only new specification is contract_after_failure, which specifies the number of unsuccessful cycles which must occur with a specific delta prior to contraction of the delta.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

See Also

These keywords may also be of interest:

- coliny_beta
7.2 method

- coliny_direct
- coliny_pattern_search
- coliny_cobyla
- coliny_ea

7.2.36.1 contract_after_failure

- Keywords Area
- method
- coliny_solis_wets
- contract_after_failure

The number of unsuccessful cycles prior to contraction.

Specification

Alias: none
Argument(s): INTEGER
Default: 4+number of variables

Description

In particular, coliny_solis_wets supports dynamic rescaling of the step length, and dynamic rescaling of the constraint penalty. The only new specification is contract_after_failure, which specifies the number of unsuccessful cycles which must occur with a specific delta prior to contraction of the delta.

7.2.36.2 no_expansion

- Keywords Area
- method
- coliny_solis_wets
- no_expansion

Don’t allow expansion of the search pattern

Specification

Alias: none
Argument(s): none
Default: algorithm may expand pattern size
In general, pattern search methods can expand and contract their step lengths. SCOLIB pattern search methods contract the step length by the value $\text{contraction\_factor}$, and they expand the step length by the value $(1/\text{contraction\_factor})$. The $\text{expand\_after\_success}$ control specifies how many successful objective function improvements must occur with a specific step length prior to expansion of the step length, whereas the $\text{no\_expansion}$ flag instructs the algorithm to forgo pattern expansion altogether.

### 7.2.36.3 expand\_after\_success

- **Keywords Area**
- method
- coliny\_solis\_wets
- expand\_after\_success

Set the factor by which a search pattern can be expanded

#### Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 5

#### Description

In general, pattern search methods can expand and contract their step lengths. SCOLIB pattern search methods contract the step length by the value $\text{contraction\_factor}$, and they expand the step length by the value $(1/\text{contraction\_factor})$. The $\text{expand\_after\_success}$ control specifies how many successful objective function improvements must occur with a specific step length prior to expansion of the step length, whereas the $\text{no\_expansion}$ flag instructs the algorithm to forgo pattern expansion altogether.

### 7.2.36.4 constant\_penalty

- **Keywords Area**
- method
- coliny\_solis\_wets
- constant\_penalty

Use a simple weighted penalty to manage feasibility

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** algorithm dynamically adapts the constraint penalty
7.2 method

Description

Finally, constraint infeasibility can be managed in a somewhat more sophisticated manner than the simple weighted penalty function. If the constant_penalty specification is used, then the simple weighted penalty scheme described above is used. Otherwise, the constraint penalty is adapted to the value constraint_penalty/L, where L is the the smallest step length used so far.

7.2.36.5 contraction_factor

- Keywords Area
- method
- coliny_solis_wets
- contraction_factor

Amount by which step length is rescaled

Specification

Alias: none
Argument(s): REAL
Default: 0.5

Description

For pattern search methods, contraction_factor specifies the amount by which step length is rescaled after unsuccessful iterates, must be strictly between 0 and 1.

For methods that can expand the step length, the expansion is 1/contraction_factor

7.2.36.6 constraint_penalty

- Keywords Area
- method
- coliny_solis_wets
- constraint_penalty

Multiplier for the penalty function

Specification

Alias: none
Argument(s): REAL
Default: 1.0
Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds \texttt{constraint_penalty} times the sum of squares of the constraint violations to the objective function. The default value of \texttt{constraint_penalty} is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.

### 7.2.36.7 initial_delta

- **Keywords Area**
- **method**
- **coliny_solis_wets**
- **initial_delta**

Initial step size for derivative-free optimizers

#### Specification

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0 (COBYLA), 0.1*range (PS, SW)

#### Description

The \texttt{initial_delta} keyword defines the size of the first search step in derivative-free optimization methods, specifically \texttt{asynch_pattern_search}, \texttt{coliny_cobyla}, \texttt{coliny_pattern_search}, \texttt{coliny_solis_wets}, and \texttt{mesh_adaptive_search}. It is applied in an absolute sense to all search directions.

#### Default Behavior

The default value is 1.0.

#### Usage Tips

It is recommended that \texttt{initial_delta} be the approximate distance from the initial point to the solution. If this distance is unknown, it is advisable to err on the side of choosing an \texttt{initial_delta} that is too large or to not specify it. Relative application of \texttt{initial_delta} is not available unless the user scales the problem accordingly.

#### Examples

Three example method input blocks appear below.  
For \texttt{asynch_pattern_search}:

```plaintext
method
 asynch_pattern_search
  initial_delta = .5
  contraction_factor = 0.25
  merit_function merit1_smooth
  smoothing_factor = 1.0
  constraint_tolerance = 1.e-6
```

For \texttt{coliny_pattern_search}:

```plaintext
method
 coliny_pattern_search
```
7.2 method

\begin{verbatim}
initial_delta = .2
variable_tolerance = 1.e-4
max_iterations 100
solution_accuracy = 1.e-6
seed = 1234
max_function_evaluations = 1000
\end{verbatim}

For \texttt{mesh_adaptive_search}

\begin{verbatim}
method
  mesh_adaptive_search
    initial_delta = 2.0
    seed = 1234
\end{verbatim}

7.2.36.8 \texttt{variable_tolerance}

- Keywords Area
- method
- \texttt{coliny_solis_wets}
- \texttt{variable_tolerance}

Step length-based stopping criteria for derivative-free optimizers

**Specification**

Alias: none

Argument(s): REAL

Default: 1.0e-4 (COBYLA), 1.0e-5 (PS), 1.0e-6 (SW)

**Description**

The \texttt{variable_tolerance} keyword defines the minimum step length allowed by the optimizer and is used to determine convergence. It is applicable to \texttt{asynch_pattern_search}, \texttt{coliny_cobyla}, \texttt{coliny_pattern_search}, \texttt{coliny_solis_wets}, and \texttt{mesh_adaptive_search}.

**Default Behavior**

The default value varies according to method as follows:

- \texttt{asynch_pattern_search}: 1.0e-2
- \texttt{coliny_cobyla}: 1.0e-4
- \texttt{coliny_pattern_search}: 1.0e-5
- \texttt{coliny_solis_wets}: 1.0e-6
- \texttt{mesh_adaptive_search}: 1.0e-6

**Usage Tips**

It is recommended that \texttt{variable_tolerance} be set to a value for which changes of that scale in parameter values cause negligible changes in the objective function.
Examples

Three example method input blocks appear below.

For \texttt{asynch\_pattern\_search}:

\begin{verbatim}
method asynch_pattern_search
  contraction_factor = 0.25
  variable_tolerance = 1.e-4
  solution_target = 1.e-6
  max_function_evaluations 500
  constraint_tolerance 1.e-6
\end{verbatim}

For \texttt{coliny\_pattern\_search}:

\begin{verbatim}
method coliny_pattern_search
  initial_delta = .2
  variable_tolerance = 1.e-4
  max_iterations 100
  solution_accuracy = 1.e-6
  seed = 1234
  max_function_evaluations = 1000
\end{verbatim}

For \texttt{mesh\_adaptive\_search}

\begin{verbatim}
method mesh_adaptive_search
  variable_tolerance = 0.01
  seed = 1234
\end{verbatim}

7.2.36.9 \texttt{solution\_target}

- Keywords Area
- method
- \texttt{coliny\_solis\_wets}
- \texttt{solution\_target}

Stopping criteria based on objective function value

Specification

\textbf{Alias:} solution\_accuracy

\textbf{Argument(s):} REAL

\textbf{Default:} -DBL\_MAX

\textbf{Description}

\texttt{solution\_target} is a termination criterion. The algorithm will terminate when the function value falls below \texttt{solution\_target}.

7.2.36.10 \texttt{seed}

- Keywords Area
- method
7.2 method

- `coliny_solis_wets`
- `seed`

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.36.11 show_misc_options

- `Keywords Area`
- `method`
- `coliny_solis_wets`
- `show_misc_options`

Show algorithm parameters not exposed in Dakota input

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no dump of specification options
All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

### 7.2.36.12 misc_options

- **Keywords Area**
- **method**
- **coliny_solis_wets**
- **misc_options**

Set method options not available through Dakota spec

### Specification

**Alias:** none  
**Argument(s):** STRINGLIST  
**Default:** no misc options

### Description

All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

### 7.2.36.13 max_iterations

- **Keywords Area**
- **method**
- **coliny_solis_wets**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed $\text{max}\_\text{iterations}$ iterations. In cases where there are multiple nested iterations to be controlled, $\text{max}\_\text{iterations}$ generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.36.14 convergence_tolerance

- Keywords Area
- method
- coliny_solis_wets
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4
Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a **relative convergence tolerance** for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.36.15 max_function_evaluations

- Keywords Area
- method
- coliny_solis_wets
- `max_function_evaluations`

Number of function evaluations allowed for optimizers

**Topics**

This keyword is related to the topics:

- `method_independent_controls`

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed `max_function_evaluations` evaluations. See also `max_iterations`. 
Some optimizers (e.g., ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

**Default Behavior**

Default value is 1000.

7.2.36.16 scaling

- Keywords Area
- method
- coliny_solis_wets
- scaling

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *scales* specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0e10\times\text{DBL_MIN}\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```
7.2 method

- coliny_solis_wets
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.37  coliny_cobyla

• Keywords Area

• method

• coliny_cobyla

Constrained Optimization BY Linear Approximations (COBYLA)

Topics

This keyword is related to the topics:

• package_scolib

• package_coliny

• local_optimization_methods

• constrained

Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>initial_delta</td>
<td></td>
<td>Reasonable initial changes to optimization variables</td>
</tr>
<tr>
<td>Optional</td>
<td>variable_tolerance</td>
<td></td>
<td>Required or expected accuracy in optimization variables</td>
</tr>
<tr>
<td>Optional</td>
<td>solution_target</td>
<td></td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>show_misc_options</td>
<td></td>
<td>Show algorithm parameters not exposed in Dakota input</td>
</tr>
<tr>
<td>Optional</td>
<td>misc_options</td>
<td></td>
<td>Set method options not available through Dakota spec</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
</tbody>
</table>
Description

The Constrained Optimization BY Linear Approximations (COBYLA) algorithm is an extension to the Nelder-Mead simplex algorithm for handling general linear/nonlinear constraints and is invoked using the coliny_cobyla group specification. The COBYLA algorithm employs linear approximations to the objective and constraint functions, the approximations being formed by linear interpolation at N+1 points in the space of the variables. We regard these interpolation points as vertices of a simplex. The step length parameter controls the size of the simplex and it is reduced automatically from initial_delta to variable_tolerance. One advantage that COBYLA has over many of its competitors is that it treats each constraint individually when calculating a change to the variables, instead of lumping the constraints together into a single penalty function.

See the page package_scolib for important information regarding all SCOLIB methods

coliny_cobyla is inherently serial.

Stopping Criteria

COBYLA currently only supports termination based on

- max_function_evaluations
- solution_target

Other method-independent stopping criteria (max_iterations and convergence_tolerance) will be ignored if set.

Known Bugs

The implementation of the coliny_cobyla optimization method is such that the best function value is not always returned to Dakota for reporting. The user is advised to look through the Dakota screen output or the tabular output file (if generated) to confirm what the best function value and corresponding parameter values are.

The coliny_cobyla optimization method does not always respect bound constraints when scaling is turned on.

Neither bug will be fixed, as maintaining third-party source code (such as COBYLA) is outside of the Dakota project scope.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

See Also

These keywords may also be of interest:

- coliny_beta
- coliny_direct
- coliny_pattern_search
7.2 method

• coliny_ea
• coliny_solis_wets

7.2.37.1 initial_delta

• Keywords Area
• method
• coliny_cobyla
• initial_delta

Reasonable initial changes to optimization variables

Specification

Alias: none
Argument(s): REAL
Default: 1.0 (COBYLA), 0.1×\text{range} (PS, SW)

Description

initial_delta for COBYLA should be set to be a value that is reasonable for initial changes to the optimization variables. It represents a distance from the initial simplex within which linear approximation subproblems can be trusted to be sufficiently representative of the true problem. It is analogous to the initial trust-region size used in trust-region methods.

Default Behavior
The default value is 1.0.

7.2.37.2 variable_tolerance

• Keywords Area
• method
• coliny_cobyla
• variable_tolerance

Required or expected accuracy in optimization variables.

Specification

Alias: none
Argument(s): REAL
Default: 1.0e-4 (COBYLA), 1.0e-5 (PS), 1.0e-6 (SW)
Description

Unlike its use as a stopping criteria in other methods, `variable_tolerance` is used to communicate the accuracy of the optimization variables to COBYLA. It represents the minimum distance from the simplex within which linear approximation subproblems can be trusted to be sufficiently representative of the true problem. It is analogous to the minimum trust-region size used in trust-region methods. Note that, per COBYLA documentation, the level of accuracy is not guaranteed.

**Default Behavior**

The default value is 0.0001.

**Additional Discussion**

While `variable_tolerance` is not a stopping criteria for COBYLA, we note that it may have a side effect regarding convergence. In particular, if lower accuracy is required of the optimization variables, it may stop sooner than if higher accuracy is required.

### 7.2.37.3 solution_target

- **Keywords Area**
- **method**
- **coliny_cobyla**
- **solution_target**

Stopping criteria based on objective function value

**Specification**

- **Alias:** `solution_accuracy`
- **Argument(s):** `REAL`
- **Default:** `-DBL_MAX`

**Description**

`solution_target` is a termination criterion. The algorithm will terminate when the function value falls below `solution_target`.

### 7.2.37.4 seed

- **Keywords Area**
- **method**
- **coliny_cobyla**
- **seed**

Seed of the random number generator
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior
If not specified, the seed is randomly generated.

Expected Output
If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips
If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.37.5 show_misc_options

- Keywords Area
- method
- coliny_cobyla
- show_misc_options

Show algorithm parameters not exposed in Dakota input

Specification

Alias: none
Argument(s): none
Default: no dump of specification options

Description

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The
**misc_options**

Optional specification provides a means for inputing additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

### 7.2.37.6 misc_options

- **Keywords Area**
- **method**
- **coliny_cobyla**
- **misc_options**

Set method options not available through Dakota spec

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** no misc options

**Description**

All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputing additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

### 7.2.37.7 max_iterations

- **Keywords Area**
- **method**
- **coliny_cobyla**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- **method_independent_controls**
7.2 method

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_{reliability, interval_est, evidence} / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max_function_evaluations}

7.2.37.8 convergence_tolerance

- Keywords Area
- method
- coliny_cobyla
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- \texttt{method_independent_controls}

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The \texttt{convergence_tolerance} specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.37.9 max_function_evaluations

- Keywords Area
- method
- coliny_cobyla
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.
7.2.37.10 scaling

- Keywords Area
- method
- coliny_cobyla
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.
2. **auto** - automatic scaling

First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

- two-sided bounds scaled into the interval [0,1];
- one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
- no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling

First, any characteristic values from the optional *scales* specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

### Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.37.11 **model_pointer**

- **Keywords Area**
- **method**
- **coliny_cobyla**
- **model_pointer**

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
```
variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = ’x1’ ’x2’

interface
  id_interface = ’I1’
  system async evaluation_concurrency = 5
  analysis_driver = ’text_book’

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.38 coliny_direct

  • Keywords Area
  • method
  • coliny_direct

Dividing RECTangles method

Topics

This keyword is related to the topics:

  • package_scolib
  • package_coliny
  • global_optimization_methods

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>division</td>
<td>division</td>
<td>Determine how rectangles are subdivided</td>
</tr>
<tr>
<td>Optional</td>
<td>global_balance_-parameter</td>
<td>global_balance_-parameter</td>
<td>Tolerance for whether a subregion is worth dividing</td>
</tr>
<tr>
<td>Optional</td>
<td>local_balance_-parameter</td>
<td>local_balance_-parameter</td>
<td>Tolerance for whether a subregion is worth dividing</td>
</tr>
</tbody>
</table>
7.2 method

<table>
<thead>
<tr>
<th>Optional</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>max_boxsize_limit</td>
<td>Stopping Criterion based on longest edge of hyperrectangle</td>
</tr>
<tr>
<td>min_boxsize_limit</td>
<td>Stopping Criterion based on shortest edge of hyperrectangle</td>
</tr>
<tr>
<td>constraint_penalty</td>
<td>Multiplier for the penalty function</td>
</tr>
<tr>
<td>solution_target</td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>show_misc_options</td>
<td>Show algorithm parameters not exposed in Dakota input</td>
</tr>
<tr>
<td>misc_options</td>
<td>Set method options not available through Dakota spec</td>
</tr>
<tr>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>max_function_-evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

Description

The Dividing RECTangles (DIRECT) optimization algorithm is a derivative free global optimization method that balances local search in promising regions of the design space with global search in unexplored regions. As shown in Figure 5.1, DIRECT adaptively subdivides the space of feasible design points so as to guarantee that iterates are generated in the neighborhood of a global minimum in finitely many iterations.

DIRECT" image latex direct1.eps "Design space partitioning with DIRECT" width=10cm
In practice, DIRECT has proven an effective heuristic for engineering design applications, for which it is able to quickly identify candidate solutions that can be further refined with fast local optimizers.

**See the page package_scolib for important information regarding all SCOLIB methods**

The DIRECT algorithm supports concurrency up to twice the number of variables being optimized.

DIRECT uses the `solution_target`, `constraint_penalty` and `show_misc_options` specifications that are described in package_scolib. Note, however, that DIRECT uses a fixed penalty value for constraint violations (i.e. it is not dynamically adapted as is done in `coliny_pattern_search`).

**Search Parameters**

The `global_balance_parameter` controls how much global search is performed by only allowing a subregion to be subdivided if the size of the subregion divided by the size of the largest subregion is at least `global_balance_parameter`. Intuitively, this forces large subregions to be subdivided before the smallest subregions are refined. The `local_balance_parameter` provides a tolerance for estimating whether the smallest subregion can provide a sufficient decrease to be worth subdividing; the default value is a small value that is suitable for most applications.

**Stopping Criteria**

DIRECT can be terminated with:

- `max_function_evaluations`
- `max_iterations`
- `convergence_tolerance`
- `solution_target`
- `max_boxsize_limit`
- `min_boxsize_limit` - most effective in practice

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions` are specified)
- Best Nonlinear Constraints
- Calibration (when `calibration_terms` are specified)

**See Also**

These keywords may also be of interest:

- `coliny_beta`
- `coliny_pattern_search`
- `coliny_cobyla`
- `coliny_ea`
- `coliny_solis_wets`
- `ncsu_direct`
7.2.38.1  division

- Keywords Area
- method
- coliny_direct
- division

Determine how rectangles are subdivided

Specification

Alias: none
Argument(s): none
Default: major_dimension

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Divide Along (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>major_dimension</td>
<td>(default) Longest edge of subregion is subdivided</td>
</tr>
<tr>
<td></td>
<td></td>
<td>all_dimensions</td>
<td>All dimensions are simultaneously subdivided</td>
</tr>
</tbody>
</table>

Description

The division specification determines how DIRECT subdivides each subregion of the search space.

If division is set to major_dimension, then the dimension representing the longest edge of the subregion is subdivided (this is the default). If division is set to all_dimensions, then all dimensions are simultaneously subdivided.

7.2.38.1.1  major_dimension

- Keywords Area
- method
- coliny_direct
- division
- major_dimension

(default) Longest edge of subregion is subdivided

Specification

Alias: none
Argument(s): none
Description

Longest edge of subregion is subdivided

7.2.38.1.2 all_dimensions

• Keywords Area
• method
• coliny_direct
• division
• all_dimensions

All dimensions are simultaneously subdivided

Specification

Alias: none
Argument(s): none

Description

All dimensions are simultaneously subdivided

7.2.38.2 global_balance_parameter

• Keywords Area
• method
• coliny_direct
• global_balance_parameter

Tolerance for whether a subregion is worth dividing

Specification

Alias: none
Argument(s): REAL
Default: 0.0

Description

The global_balance_parameter controls how much global search is performed by only allowing a subregion to be subdivided if the size of the subregion divided by the size of the largest subregion is at least global_balance_parameter. Intuitively, this forces large subregions to be subdivided before the smallest subregions are refined.
7.2 method

7.2.38.3 local_balance_parameter

• Keywords Area

• method

• coliny_direct

• local_balance_parameter

Tolerance for whether a subregion is worth dividing

Specification

Alias: none

Argument(s): REAL

Default: 1.e-8

Description

See parent page. The local_balance_parameter provides a tolerance for estimating whether the smallest subregion can provide a sufficient decrease to be worth subdividing; the default value is a small value that is suitable for most applications.

7.2.38.4 max_boxsize_limit

• Keywords Area

• method

• coliny_direct

• max_boxsize_limit

Stopping Criterion based on longest edge of hyperrectangle

Specification

Alias: none

Argument(s): REAL

Default: 0.0

Description

Each subregion considered by DIRECT has a size, which corresponds to the longest diagonal of the subregion. max_boxsize_limit specification terminates DIRECT if the size of the largest subregion falls below this threshold.
7.2.38.5 min_boxsize_limit

- Keywords Area
- method
- coliny_direct
- min_boxsize_limit

Stopping Criterion based on shortest edge of hyperrectangle

Specification

Alias: none

Argument(s): REAL

Default: 1.0e-4

Description

min_boxsize_limit is a setting that terminates the optimization when the measure of a hyperrectangle S with \( f(c(S)) = f_{\text{min}} \) is less than \( \text{min\_boxsize\_limit} \).

Each subregion considered by DIRECT has a size, which corresponds to the longest diagonal of the subregion. \( \text{min\_boxsize\_limit} \) specification terminates DIRECT if the size of the smallest subregion falls below this threshold.

In practice, this specification is likely to be more effective at limiting DIRECT's search.

7.2.38.6 constraint_penalty

- Keywords Area
- method
- coliny_direct
- constraint_penalty

Multiplier for the penalty function

Specification

Alias: none

Argument(s): REAL

Default: 1000.0

Description

Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds \( \text{constraint\_penalty} \) times the sum of squares of the constraint violations to the objective function. The default value of \( \text{constraint\_penalty} \) is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.
7.2 method

7.2.38.7 solution_target

- Keywords Area
- method
- coliny_direct
- solution_target

Stopping criteria based on objective function value

Specification

Alias: solution_accuracy
Argument(s): REAL
Default: -DBL_MAX

Description

solution_target is a termination criterion. The algorithm will terminate when the function value falls below solution_target.

7.2.38.8 seed

- Keywords Area
- method
- coliny_direct
- seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips
If a stochastic study was run without \textit{seed} specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

\textbf{Examples}

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
\end{verbatim}

\textbf{7.2.38.9 show\_misc\_options}

\begin{itemize}
  \item Keywords Area
  \item method
  \item coliny\_direct
  \item show\_misc\_options
\end{itemize}

Show algorithm parameters not exposed in Dakota input

\textbf{Specification}

\textbf{Alias:} none

\textbf{Argument(s):} none

\textbf{Default:} no dump of specification options

\textbf{Description}

All SCOLIB methods support the \textit{show\_misc\_options} optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The \textit{misc\_options} optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

\textbf{7.2.38.10 misc\_options}

\begin{itemize}
  \item Keywords Area
  \item method
  \item coliny\_direct
  \item misc\_options
\end{itemize}

Set method options not available through Dakota spec
7.2 method

Specification

Alias: none
Argument(s): STRINGLIST
Default: no misc options

Description

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The misc_options optional specification provides a means for inputing additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

7.2.38.11 max_iterations

- Keywords Area
- method
- coliny_direct
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25\(\times n\))

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.38.12 convergence_tolerance

- Keywords Area
- method
- coliny_direct
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

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7.2.38.13 max_function_evaluations

* Keywords Area
  * method
  * coliny_direct
  * max_function_evaluations

Number of function evaluations allowed for optimizers

**Topics**

This keyword is related to the topics:

  * method_independent_controls

**Specification**

*Alias*: none

*Argument(s)*: INTEGER

*Default*: 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

**Default Behavior**

Default value is 1000.

7.2.38.14 scaling

* Keywords Area
  * method
  * coliny_direct
  * scaling

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

  * method_independent_controls
Specification

**Alias:** none

**Argument(s):** none

**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `$scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval \([0,1]\);
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into \([0,1]\).

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.
Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0 \times 10^{-10} \times \text{DBL_MIN}\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.38.15 model_pointer

- **Keywords Area**
- **method**
- **coliny_direct**
- **model_pointer**

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- **block_pointer**

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)
Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling, 
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6 
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians
7.2.39 coliny_ea

- **Keywords Area**
- **method**
- **coliny_ea**

Evolutionary Algorithm

**Topics**

This keyword is related to the topics:

- **package_scolib**
- **package_coliny**
- **global_optimization_methods**

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>population_size</td>
<td></td>
<td>Set the population size</td>
</tr>
<tr>
<td>Optional</td>
<td>initialization_type</td>
<td></td>
<td>Specify how to initialize</td>
</tr>
<tr>
<td>Optional</td>
<td>fitness_type</td>
<td></td>
<td>Select fitness type</td>
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<td></td>
<td>Select a replacement type</td>
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<td>crossover_rate</td>
<td></td>
<td>Specify the probability</td>
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<tr>
<td>Optional</td>
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<td></td>
<td>Select a crossover type</td>
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<tr>
<td>Optional</td>
<td>mutation_rate</td>
<td></td>
<td>Set probability of a mutation</td>
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<td>Optional</td>
<td>mutation_type</td>
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<td>Optional</td>
<td>constraint_penalty</td>
<td></td>
<td>Multiplier for the penalty function</td>
</tr>
<tr>
<td>Optional</td>
<td>solution_target</td>
<td>Stopping criteria based on objective function value</td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>-----------------</td>
<td>---------------------------------------------------</td>
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</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>show_mis_correct</td>
<td>Show algorithm parameters not exposed in Dakota input</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>misc_options</td>
<td>Set method options not available through Dakota spec</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
<td></td>
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<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
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<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
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</tbody>
</table>

### Description

**Evolutionary Algorithm**

See the page `package_scolib` for important information regarding all SCOLIB methods.

`coliny_pattern_search` supports concurrency up to the size of the population.

The random `seed` control provides a mechanism for making a stochastic optimization repeatable. That is, the use of the same random seed in identical studies will generate identical results. The `population_size` control specifies how many individuals will comprise the EA’s population.

The `initialization_type` defines the type of initialization for the population of the EA. There are three types: `simple_random`, `unique_random`, and `flat_file`.  `simple_random` creates initial solutions with random variable values according to a uniform random number distribution. It gives no consideration to any previously generated designs. The number of designs is specified by the `population_size`.  `unique_random` is the same as `simple_random`, except that when a new solution is generated, it is checked against the rest of the solutions. If it duplicates any of them, it is rejected.  `flat_file` allows the initial population to be read from a flat file. If `flat_file` is specified, a file name must be given.
The **fitness_type** controls how strongly differences in "fitness" (i.e., the objective function) are weighted in the process of selecting "parents" for crossover:

- the **linear_rank** setting uses a linear scaling of probability of selection based on the rank order of each individual’s objective function within the population

- the **merit_function** setting uses a proportional scaling of probability of selection based on the relative value of each individual’s objective function within the population

The **replacement_type** controls how current populations and newly generated individuals are combined to create a new population. Each of the **replacement_type** selections accepts an integer value, which is referred to below as the **replacement_size**.

- The **random** setting creates a new population using (a) **replacement_size** randomly selected individuals from the current population, and (b) **population_size** · **replacement_size** individuals randomly selected from among the newly generated individuals (the number of which is optionally specified using **new_solutions_generated**) that are created for each generation (using the selection, crossover, and mutation procedures).

- The **chc** setting creates a new population using (a) the **replacement_size** best individuals from the **combination** of the current population and the newly generated individuals, and (b) **population_size** · **replacement_size** individuals randomly selected from among the remaining individuals in this combined pool. The chc setting is the preferred selection for many engineering problems.

- The **elitist** (default) setting creates a new population using (a) the **replacement_size** best individuals from the current population, (b) and **population_size** · **replacement_size** individuals randomly selected from the newly generated individuals. It is possible in this case to lose a good solution from the newly generated individuals if it is not randomly selected for replacement; however, the default **new_solutions_generated** value is set such that the entire set of newly generated individuals will be selected for replacement.

Note that **new_solutions_generated** is not recognized by Dakota as a valid keyword unless **replacement_type** has been specified.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the **hdf5** keyword, this method writes the following results to HDF5:

- **Best Parameters**

- **Best Objective Functions** (when **objective_functions**) are specified)

- **Best Nonlinear Constraints**

- **Calibration** (when **calibration_terms** are specified)

**Theory**

The basic steps of an evolutionary algorithm are depicted in Figure 5.2.
They can be enumerated as follows:

1. Select an initial population randomly and perform function evaluations on these individuals
2. Perform selection for parents based on relative fitness
3. Apply crossover and mutation to generate new_solutions_generated new individuals from the selected parents
   - Apply crossover with a fixed probability from two selected parents
   - If crossover is applied, apply mutation to the newly generated individual with a fixed probability
   - If crossover is not applied, apply mutation with a fixed probability to a single selected parent
4. Perform function evaluations on the new individuals
5. Perform replacement to determine the new population
6. Return to step 2 and continue the algorithm until convergence criteria are satisfied or iteration limits are exceeded

See Also

These keywords may also be of interest:

- coliny_beta
- coliny_direct
- coliny_pattern_search
- coliny_cobyla
- coliny_solis_wets

7.2.39.1 population_size

- Keywords Area
- method
7.2 method

- `coliny_ea`
- `population_size`

Set the population size

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 50

**Description**

The number of designs in the population is specified by the `population_size`.

7.2.39.2 `initialization_type`

- **Keywords Area**  
  - `method`  
  - `coliny_ea`  
  - `initialization_type`

Specify how to initialize the population

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** unique_random

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Initialization Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
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<td></td>
<td><code>simple_random</code></td>
<td>Create random initial solutions</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>unique_random</code></td>
<td>Create random initial solutions, but enforce uniqueness (default)</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>flat_file</code></td>
<td>Read initial solutions from file</td>
</tr>
</tbody>
</table>

**Description**

The `initialization_type` defines how the initial population is created for the GA. There are three types:

1. `simple_random`
2. `unique_random` (default)
3. flat_file

Setting the size for the flat_file initializer has the effect of requiring a minimum number of designs to create. If this minimum number has not been created once the files are all read, the rest are created using the unique_random initializer and then the simple_random initializer if necessary.

7.2.39.2.1 simple_random

- Keywords Area
- method
- coliny_ea
- initialization_type
- simple_random

Create random initial solutions

Specification

Alias: none
Argument(s): none

Description

simple_random creates initial solutions with random variable values according to a uniform random number distribution. It gives no consideration to any previously generated designs.

7.2.39.2.2 unique_random

- Keywords Area
- method
- coliny_ea
- initialization_type
- unique_random

Create random initial solutions, but enforce uniqueness (default)

Specification

Alias: none
Argument(s): none

Description

unique_random is the same as simple_random, except that when a new solution is generated, it is checked against the rest of the solutions. If it duplicates any of them, it is rejected.
7.2.39.3 flat_file

- Keywords Area
- method
- coliny_ea
- initialization_type
- flat_file

Read initial solutions from file

Specification

Alias: none
Argument(s): STRING

Description

flat_file allows the initial population to be read from a flat file. If flat_file is specified, a file name must be given.

7.2.39.3 fitness_type

- Keywords Area
- method
- coliny_ea
- fitness_type

Select fitness type

Specification

Alias: none
Argument(s): none
Default: linear_rank

Child Keywords:

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<th>Dakota Keyword</th>
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<tr>
<td></td>
<td>merit_function</td>
<td>linear_rank</td>
<td>Balance goals of reducing objective function and satisfying constraints</td>
</tr>
</tbody>
</table>
The *fitness_type* controls how strongly differences in "fitness" (i.e., the objective function) are weighted in the process of selecting "parents" for crossover. It has two options, *linear_rank* and *merit_function*.

### 7.2.39.3.1 linear_rank

- **Keywords Area**
- **method**
- **coliny_ea**
- **fitness_type**
- **linear_rank**

Set selection scaling

### Specification

**Alias:** none

**Argument(s):** none

### Description

The *linear_rank* setting uses a linear scaling of probability of selection based on the rank order of each individual's objective function within the population.

### 7.2.39.3.2 merit_function

- **Keywords Area**
- **method**
- **coliny_ea**
- **fitness_type**
- **merit_function**

Balance goals of reducing objective function and satisfying constraints

### Specification

**Alias:** none

**Argument(s):** none

### Description

A *merit_function* is a function in constrained optimization that attempts to provide joint progress toward reducing the objective function and satisfying the constraints.
7.2 method

7.2.39.4 replacement_type

- Keywords Area
- method
- coliny_ea
- replacement_type

Select a replacement type for SCOLIB evolutionary algorithm (coliny_ea)

Specification

Alias: none
Argument(s): none
Default: elitist=1
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Replacement Type (Group 1)</td>
<td>random</td>
<td>Create new population randomly</td>
</tr>
<tr>
<td>chc</td>
<td></td>
<td></td>
<td>Create new population using replacement</td>
</tr>
<tr>
<td>elitist</td>
<td></td>
<td></td>
<td>Use the best designs to form a new population</td>
</tr>
<tr>
<td>Optional</td>
<td>new_solutions_generated</td>
<td></td>
<td>Replace population with individuals chosen from population</td>
</tr>
</tbody>
</table>

Description

The replacement_type controls how current populations and newly generated individuals are combined to create a new population. Each of the replacement_type selections accepts an associated integer value, which is specified by the replacement_size:

The random setting creates a new population using (a) replacement_size randomly selected individuals from the current population, and (b) population_size - replacement_size individuals randomly selected from among the newly generated individuals (the number of which is optionally specified using new_solutions_generated) that are created for each generation (using the selection, crossover, and mutation procedures).

The chc setting creates a new population using (a) the replacement_size best individuals from the combination of the current population and the newly generated individuals, and (b) population_size - replacement_size individuals randomly selected from among the remaining individuals in this combined pool. The chc setting is the preferred selection for many engineering problems.

The elitist (default) setting creates a new population using (a) the replacement_size best individuals from the current population, (b) and population_size - replacement_size individuals randomly selected from the newly generated individuals. It is possible in this case to lose a good solution from the newly generated individuals if it is not randomly selected for replacement; however, the default new_solutions_generated value is set such that the entire set of newly generated individuals will be selected for replacement.

Note that new_solutions_generated is not recognized by Dakota as a valid keyword unless replacement_type has been specified.
7.2.39.4.1 random

- Keywords Area
- method
- coliny_ea
- replacement_type
- random

Create new population randomly

Specification

Alias: none
Argument(s): INTEGER

Description

The replacement_type controls how current populations and newly generated individuals are combined to create a new population. Each of the replacement_type selections accepts an integer value, which is referred as the replacement_size:

The random setting creates a new population using:

- replacement_size randomly selected individuals from the current population, and

- population_size - replacement_size individuals randomly selected from among the newly generated individuals (the number of which is optionally specified using new_solutions_generated) that are created for each generation (using the selection, crossover, and mutation procedures).

7.2.39.4.2 chc

- Keywords Area
- method
- coliny_ea
- replacement_type
- chc

Create new population using replacement

Specification

Alias: none
Argument(s): INTEGER
Description

The replacement_type controls how current populations and newly generated individuals are combined to create a new population. Each of the replacement_type selections accepts an integer value, which is referred to as the replacement_size:

The chc setting creates a new population using (a) the replacement_size best individuals from the combination of the current population and the newly generated individuals, and (b) population_size - replacement_size individuals randomly selected from among the remaining individuals in this combined pool. The chc setting is the preferred selection for many engineering problems.

7.2.39.4.3 elitist

- Keywords Area
- method
- coliny_ea
- replacement_type
- elitist

Use the best designs to form a new population

Specification

Alias: none
Argument(s): INTEGER

Description

The elitist (default) setting creates a new population using (a) the replacement_size best individuals from the current population, (b) and population_size - replacement_size individuals randomly selected from the newly generated individuals. It is possible in this case to lose a good solution from the newly generated individuals if it is not randomly selected for replacement; however, the default new_solutions_generated value is set such that the entire set of newly generated individuals will be selected for replacement.

7.2.39.4.4 new_solutions_generated

- Keywords Area
- method
- coliny_ea
- replacement_type
- new_solutions_generated

Replace population with individuals chosen from population

Specification

Alias: none
Argument(s): INTEGER
Default: population_size - replacement_size
Description

- The random setting creates a new population using (a) \textit{replacement\_size} randomly selected individuals from the current population, and (b) \textit{population\_size} - \textit{replacement\_size} individuals randomly selected from among the newly generated individuals (the number of which is optionally specified using \textit{new\_solutions\_generated}) that are created for each generation (using the selection, crossover, and mutation procedures).

7.2.39.5 \textit{crossover\_rate}

\begin{itemize}
  \item \textbf{Keywords Area}
  \item \textbf{method}
  \item \textbf{coliny\_ea}
  \item \textbf{crossover\_rate}
\end{itemize}

Specify the probability of a crossover event

Specification

\textbf{Alias:} none

\textbf{Argument(s):} REAL

\textbf{Default:} 0.8

Description

The \textit{crossover\_type} controls what approach is employed for combining parent genetic information to create offspring, and the \textit{crossover\_rate} specifies the probability of a crossover operation being performed to generate a new offspring. The SCOLIB EA method supports three forms of crossover, \textit{two\_point}, \textit{blend}, and \textit{uniform}, which generate a new individual through combinations of two parent individuals. Two-point crossover divides each parent into three regions, where offspring are created from the combination of the middle region from one parent and the end regions from the other parent. Since the SCOLIB EA does not utilize bit representations of variable values, the crossover points only occur on coordinate boundaries, never within the bits of a particular coordinate. Uniform crossover creates offspring through random combination of coordinates from the two parents. Blend crossover generates a new individual randomly along the multidimensional vector connecting the two parents.

7.2.39.6 \textit{crossover\_type}

\begin{itemize}
  \item \textbf{Keywords Area}
  \item \textbf{method}
  \item \textbf{coliny\_ea}
  \item \textbf{crossover\_type}
\end{itemize}

Select a crossover type
Specification

Alias: none
Argument(s): none
Default: two_point
Child Keywords:
The `crossover_type` controls what approach is employed for combining parent genetic information to create offspring. The SCOLIB EA method supports three forms of crossover, `two_point`, `blend`, and `uniform`, which generate a new individual through combinations of two parent individuals.

### 7.2.39.6.1 two_point

- **Keywords Area**
- **method**
- **coliny_ea**
- **crossover_type**
- **two_point**

Combine middle of one parent with end of another

### Specification

**Alias:** none

**Argument(s):** none

### Description

Two-point crossover divides each parent into three regions, where offspring are created from the combination of the middle region from one parent and the end regions from the other parent. Since the SCOLIB EA does not utilize bit representations of variable values, the crossover points only occur on coordinate boundaries, never within the bits of a particular coordinate.

### 7.2.39.6.2 blend

- **Keywords Area**
- **method**
- **coliny_ea**
- **crossover_type**
- **blend**

Random blend of parents
7.2 method

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

`blend` crossover generates a new individual randomly along the multidimensional vector connecting the two parents.

7.2.39.6.3 uniform

- **Keywords Area**
- **method**
- **coliny_ea**
- **crossover_type**
- **uniform**

Randomly combine coordinates from parents

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Uniform crossover creates offspring through random combination of coordinates from the two parents.

7.2.39.7 mutation_rate

- **Keywords Area**
- **method**
- **coliny_ea**
- **mutation_rate**

Set probability of a mutation

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0
Description

The `mutation_rate` controls the probability of mutation being performed on an individual, both for new individuals generated by crossover (if crossover occurs) and for individuals from the existing population. It is the fraction of trial points that are mutated in a given iteration and therefore must be specified to be between 0 and 1.

7.2.39.8 mutation_type

- Keywords Area
- method
- coliny_ea
- mutation_type

Select a mutation type

Specification

Alias: none

Argument(s): none

Default: offset_normal

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Mutation Type (Group 1)</td>
<td>replace_uniform</td>
<td>Replace coordinate with randomly generated value</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_normal</td>
<td>Set mutation offset to use a normal distribution</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_cauchy</td>
<td>Use a Cauchy distribution for the mutation offset</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offset_uniform</td>
<td>Set mutation offset to use a uniform distribution</td>
</tr>
<tr>
<td>Optional</td>
<td>non_adaptive</td>
<td>Disable self-adaptive mutation</td>
<td></td>
</tr>
</tbody>
</table>

Description

The `mutation_type` controls what approach is employed in randomly modifying continuous design variables within the EA population. Each of the mutation methods generates coordinate-wise changes to individuals, usually by adding a random variable to a given coordinate value (an `offset_*` mutation), but also by replacing a given coordinate value with a random variable (a `replace_*` mutation).

Discrete design variables are always mutated using the `offset_uniform` method.

7.2.39.8.1 replace_uniform

- Keywords Area
- method
Replace coordinate with randomly generated value

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `replace_uniform` mutation type generates a replacement value for a coordinate using a uniformly distributed value over the total range for that coordinate.

**7.2.39.8.2 offset_normal**

- **Keywords Area**
  - method
  - coliny_ea
  - mutation_type
  - offset_normal

Set mutation offset to use a normal distribution

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_scale</td>
<td>Scales mutation across range of parameter</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_range</td>
<td>Set uniform offset control for discrete parameters</td>
</tr>
</tbody>
</table>

**Description**

The `offset_normal` type is an "offset" mutation that adds a 0-mean random variable with a normal uniform distribution to the existing coordinate value. The offset is limited in magnitude by `mutation_scale.`
7.2.39.8.2.1 mutation_scale

Scales mutation across range of parameter

Specification

Alias: none
Argument(s): REAL
Default: 0.1

Description

The mutation_scale specifies a scale factor which scales continuous mutation offsets; this is a fraction of the total range of each dimension, so mutation_scale is a relative value between 0 and 1.

7.2.39.8.2.2 mutation_range

Set uniform offset control for discrete parameters

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The mutation_range is used to control offset_uniform mutation used for discrete parameters. The replacement discrete value is the original value plus or minus an integer value up to mutation_range.
7.2 method

7.2.39.8.3 offset_cauchy

• Keywords Area
• method
• coliny_ea
• mutation_type
• offset_cauchy

Use a Cauchy distribution for the mutation offset

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_scale</td>
<td>Scales mutation across range of parameter</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>mutation_range</td>
<td>Set uniform offset control for discrete parameters</td>
</tr>
</tbody>
</table>

Description

The offset_cauchy type is an "offset" mutation that adds a 0-mean random variable with a cauchy distribution to the existing coordinate value. The offset is limited in magnitude by mutation_scale.

7.2.39.8.3.1 mutation_scale

• Keywords Area
• method
• coliny_ea
• mutation_type
• offset_cauchy
• mutation_scale

Scales mutation across range of parameter

Specification

Alias: none
Argument(s): REAL
Default: 0.1
Description

The mutation_scale specifies a scale factor which scales continuous mutation offsets; this is a fraction of the total range of each dimension, so mutation_scale is a relative value between 0 and 1.

7.2.39.8.3.2 mutation_range

- Keywords Area
- method
- coliny_ea
- mutation_type
- offset_cauchy
- mutation_range

Set uniform offset control for discrete parameters

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The mutation_range is used to control offset_uniform mutation used for discrete parameters. The replacement discrete value is the original value plus or minus an integer value up to mutation_range.

7.2.39.8.4 offset_uniform

- Keywords Area
- method
- coliny_ea
- mutation_type
- offset_uniform

Set mutation offset to use a uniform distribution

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

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7.2 method

| Optional | mutation_scale | Scales mutation across range of parameter Set uniform offset control for discrete parameters |
| Optional | mutation_range | |

Description

The `offset_uniform` type is an "offset" mutation that adds a 0-mean random variable with a uniform distribution to the existing coordinate value. The offset is limited in magnitude by `mutation_scale`.

For discrete design variables, `offset_uniform` is always used, and `mutation_range` controls the magnitude of the mutation.

7.2.39.8.4.1 mutation_scale

- Keywords Area
- method
- coliny_ea
- mutation_type
- offset_uniform
- mutation_scale

Scales mutation across range of parameter

Specification

Alias: none

Argument(s): REAL

Default: 0.1

Description

The `mutation_scale` specifies a scale factor which scales continuous mutation offsets; this is a fraction of the total range of each dimension, so `mutation_scale` is a relative value between 0 and 1.

7.2.39.8.4.2 mutation_range

- Keywords Area
- method
- coliny_ea
- mutation_type
- offset_uniform
- mutation_range

Set uniform offset control for discrete parameters
Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1

**Description**

The `mutation_range` is used to control `offset_uniform` mutation used for discrete parameters. The replacement discrete value is the original value plus or minus an integer value up to `mutation_range`.

7.2.39.8.5 non_adaptive

- **Keywords Area**
- **method**
- **coliny_ea**
- **mutation_type**
- **non_adaptive**

Disable self-adaptive mutation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** Adaptive mutation

**Description**

The SCOLIB EA method uses self-adaptive mutation, which modifies the mutation scale dynamically. This mechanism is borrowed from EAs like evolution strategies. The `non_adaptive` flag can be used to deactivate the self-adaptation, which may facilitate a more global search.

Note that `non_adaptive` is not recognized by Dakota as a valid keyword unless `mutation_type` has been specified.

7.2.39.9 constraint_penalty

- **Keywords Area**
- **method**
- **coliny_ea**
- **constraint_penalty**

Multiplier for the penalty function

**Specification**

**Alias:** none  
**Argument(s):** REAL
Description

Most SCOLIB optimizers treat constraints with a simple penalty scheme that adds \( \text{constraint\_penalty} \) times the sum of squares of the constraint violations to the objective function. The default value of \( \text{constraint\_penalty} \) is 1000.0, except for methods that dynamically adapt their constraint penalty, for which the default value is 1.0.

7.2.39.10 solution_target

- Keywords Area
- method
- coliny_ea
- solution_target

Stopping criteria based on objective function value

Specification

Alias: solution_accuracy
Argument(s): REAL
Default: -DBL_MAX

Description

solution_target is a termination criterion. The algorithm will terminate when the function value falls below solution_target.

7.2.39.11 seed

- Keywords Area
- method
- coliny_ea
- seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)
Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.39.12 show_misc_options

- Keywords Area
- method
- coliny_ea
- show_misc_options

Show algorithm parameters not exposed in Dakota input

Specification

Alias: none

Argument(s): none

Default: no dump of specification options

Description

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The misc_options optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.
7.2.39.13 misc_options

- Keywords Area
- method
- coliny_ea
- misc_options

Set method options not available through Dakota spec

**Specification**

*Alias*: none

*Argument(s)*: STRINGLIST

*Default*: no misc options

**Description**

All SCOLIB methods support the `show_misc_options` optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The `misc_options` optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

7.2.39.14 max_iterations

- Keywords Area
- method
- coliny_ea
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

*Alias*: none

*Argument(s)*: INTEGER

*Default*: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)
The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \textit{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \textit{max\_iterations} generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- \textit{max\_function\_evaluations}

**7.2.39.15 convergence\_tolerance**

**Keywords Area**

- \textit{method}
- \textit{coliny\_ea}
- \textit{convergence\_tolerance}

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- \textit{method\_independent\_controls}

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The \textit{convergence\_tolerance} specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a \textbf{relative convergence tolerance} for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by \textit{convergence\_tolerance}, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.39.16 max_function_evaluations

- Keywords Area
- method
- coliny_ea
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

Default Behavior

Default value is 1000.

7.2.39.17 scaling

- Keywords Area
- method
- coliny_ea
• **scaling**

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

• **method_independent_controls**

**Specification**

**Alias:** none  

**Argument(s):** none  

**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the **scaling** keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords **scale_types** and/or **scales** in the **variables** and **responses** blocks. When the **scaling** keyword is omitted from **method**, these scaling type and value specifications are ignored in the **variables** and **responses** sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a **scales** keyword, which gives characteristic values (divisors)
- a **scale_types** keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in **responses** for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. log - logarithmic scaling

First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

### Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.39.18 model_pointer

- Keywords Area
- method
- coliny_ea
- model_pointer

Identifier for model block to be used by a method

### Topics

This keyword is related to the topics:

- block_pointer
Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
7.2 method

```python
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

7.2.40 coliny_beta

- Keywords Area
- method
- coliny_beta

(Experimental) Coliny beta solver

Topics

This keyword is related to the topics:

- package_scolib
- package_coliny

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>beta_solver_name</td>
<td></td>
<td>Use an in-development SCOLIB solver</td>
</tr>
<tr>
<td>Optional</td>
<td>solution_target</td>
<td></td>
<td>Stopping criteria based on objective function value</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>show_misc_options</td>
<td></td>
<td>Show algorithm parameters not exposed in Dakota input</td>
</tr>
<tr>
<td>Optional</td>
<td>misc_options</td>
<td></td>
<td>Set method options not available through Dakota spec</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
</tr>
<tr>
<td>-------------------</td>
<td>----------------</td>
<td>---------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-</td>
<td>Stopping criterion based on objective function or statistics</td>
<td></td>
</tr>
<tr>
<td></td>
<td>tolerance</td>
<td>convergence</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
</tr>
<tr>
<td></td>
<td>evaluations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

This method keyword allows testing of experimental (beta) Coliny (Scolib) optimization solvers during software development. It is intended primarily for developer use. Additional information on Coliny solvers is available at package_scolib.

**See Also**

These keywords may also be of interest:

- coliny_direct
- coliny_pattern_search
- coliny_cobyla
- coliny_ea
- coliny_solis_wets

7.2.40.1 beta_solver_name

- Keywords Area
- method
- coliny_beta
- beta_solver_name

Use an in-development SCOLIB solver
7.2 method

Specification

Alias: none
Argument(s): STRING

Description

This is a means of accessing new methods in SCOLIB before they are exposed through the Dakota interface. Seek help from a Dakota or SCOLIB developer or a Dakota developer.

7.2.40.2 solution_target

• Keywords Area
• method
• coliny_beta
• solution_target

Stopping criteria based on objective function value

Specification

Alias: solution_accuracy
Argument(s): REAL
Default: -DBL_MAX

Description

solution_target is a termination criterion. The algorithm will terminate when the function value falls below solution_target.

7.2.40.3 seed

• Keywords Area
• method
• coliny_beta
• seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)
Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.40.4 show_misc_options

- Keywords Area
- method
- coliny_beta
- show_misc_options

Show algorithm parameters not exposed in Dakota input

Specification

Alias: none

Argument(s): none

Default: no dump of specification options

Description

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The misc_options optional specification provides a means for inputting additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification; they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.
7.2.40.5 misc_options

- Keywords Area
- method
- coliny_beta
- misc_options

Set method options not available through Dakota spec

Specification

Alias: none

Argument(s): STRINGLIST

Default: no misc options

Description

All SCOLIB methods support the show_misc_options optional specification which results in a dump of all the allowable method inputs. Note that the information provided by this command refers to optimizer parameters that are internal to SCOLIB, and which may differ from corresponding parameters used by the Dakota interface. The misc_options optional specification provides a means for inputing additional settings supported by the SCOLIB methods but which are not currently mapped through the Dakota input specification. Care must be taken in using this specification: they should only be employed by users familiar with the full range of parameter specifications available directly from SCOLIB and understand any differences that exist between those specifications and the ones available through Dakota.

7.2.40.6 max_iterations

- Keywords Area
- method
- coliny_beta
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- \( \text{max\_function\_evaluations} \)

**7.2.40.7 convergence\_tolerance**

- \( \text{Keywords Area} \)
- \( \text{method} \)
- \( \text{coliny\_beta} \)
- \( \text{convergence\_tolerance} \)

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- \( \text{method\_independent\_controls} \)

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The \( \text{convergence\_tolerance} \) specification provides a real value for controlling the termination of iteration.

For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by \( \text{convergence\_tolerance} \), then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.40.8 max_function_evaluations

This keyword is related to the following:

- **Keywords Area**
- **method**
- **coliny_beta**
- **max_function_evaluations**

#### Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1000

#### Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \( \text{max\_function\_evaluations} \) evaluations. See also \( \text{max\_iterations} \).

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before \( \text{max\_function\_evaluations} \) was exceeded.

**Default Behavior**

Default value is 1000.

### 7.2.40.9 scaling

This keyword is related to the following:

- **Keywords Area**
- **method**
- **coliny_beta**
• **scaling**

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

• **method_independent_controls**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the **scaling** keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords **scale_types** and/or **scales** in the **variables** and **responses** blocks. When the **scaling** keyword is omitted from **method**, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

• a **scales** keyword, which gives characteristic values (divisors)  
• a **scale_types** keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in **responses** for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

   • two-sided bounds scaled into the interval \([0,1]\);
   • one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
7.2 method

- no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. log - logarithmic scaling

First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.
When continuous design variables are log scaled, linear constraints are not allowed.

Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10 \times \text{DBL_MIN}$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100
```

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.40.10 model_pointer

- Keywords Area
- method
- coliny_beta
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```dakota
environment
  tabular_data
  method_pointer = 'UQ'
method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative
model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic
method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2
model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'
variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'
interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
7.2 method

```python
analysis_driver = 'text_book'
responses
response_functions = 3
no_gradients
no_hessians
```

7.2.41 nl2sol

- **Keywords Area**
- method
- nl2sol

Trust-region method for nonlinear least squares

**Topics**

This keyword is related to the topics:

- **nonlinear_least_squares**

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>function_precision</td>
<td></td>
<td>Specify the maximum precision of the analysis code responses</td>
</tr>
<tr>
<td>Optional</td>
<td>absolute_conv_tol</td>
<td></td>
<td>Absolute convergence tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td>x_conv_tol</td>
<td></td>
<td>X-convergence tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td>singular_conv_tol</td>
<td></td>
<td>Singular convergence tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td>singular_radius</td>
<td></td>
<td>Singular radius</td>
</tr>
<tr>
<td>Optional</td>
<td>false_conv_tol</td>
<td></td>
<td>False convergence tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_trust_radius</td>
<td></td>
<td>Initial trust region radius</td>
</tr>
<tr>
<td>Optional</td>
<td>covariance</td>
<td></td>
<td>Determine how the final covariance matrix is computed</td>
</tr>
<tr>
<td>Optional</td>
<td>regression_diagnostics</td>
<td>Turn on regression diagnostics</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>speculative</td>
<td>Compute speculative gradients</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

NL2SOL is available as `nl2sol` and addresses unconstrained and bound-constrained least squares problems. It uses a trust-region method (and thus can be viewed as a generalization of the Levenberg-Marquardt algorithm) and adaptively chooses between two Hessian approximations, the Gauss-Newton approximation alone and the Gauss-Newton approximation plus a quasi-Newton approximation to the rest of the Hessian. Even on small-residual problems, the latter Hessian approximation can be useful when the starting guess is far from the solution. On problems that are not over-parameterized (i.e., that do not involve more optimization variables than the data support), NL2SOL usually exhibits fast convergence.

Several internal NL2SOL convergence tolerances are adjusted in response to `function_precision`, which gives the relative precision to which responses are computed.

These tolerances may also be specified explicitly using:

- `convergence_tolerance` (NL2SOL's `rfctol`)
- `x_conv_tol` (NL2SOL's `xctol`)
- `absolute_conv_tol` (NL2SOL's `afctol`)
- `singular_conv_tol` (NL2SOL's `sctol`)
- `false_conv_tol` (NL2SOL's `xtol`)
- `initial_trust_radius` (NL2SOL's `lmax0`)

The internal NL2SOL defaults can be obtained for many of these controls by specifying the value `-1`. The internal defaults are often functions of machine epsilon (as limited by `function_precision`).
7.2 Method

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)
- Parameter Confidence Intervals (when num_experiments equals 1)

Examples

An example of nl2sol is given below, and is discussed in the User’s Manual.

Note that in this usage of calibration_terms, the driver script rosenbrock, is returning "residuals", which the nl2sol method is attempting to minimize. Another use case is to provide a data file, which Dakota will attempt to match the model responses to. See calibration_data_file. Finally, as of Dakota 6.2, the field data capability may be used with nl2sol. That is, the user can specify field simulation data and field experiment data, and Dakota will interpolate and provide the proper residuals for the calibration.

# Dakota Input File: rosen_opt_nls.in
environment
  tabular_data
tabular_data_file = 'rosen_opt_nls.dat'
method
  max_iterations = 100
  convergence_tolerance = 1e-4
nl2sol
model
  single
variables
  continuous_design = 2
  initial_point -1.2 1.0
  lower_bounds -2.0 -2.0
  upper_bounds 2.0 2.0
  descriptors 'x1' "x2"
interface
  analysis_driver = 'rosenbrock'
direct
responses
  calibration_terms = 2
  analytic_gradients
  no_hessians

Theory

NL2SOL has a variety of internal controls as described in AT&T Bell Labs CS TR 153 (http://cm.bell-labs.com/cm/cs/cstr/153.ps.gz). A number of existing Dakota controls (method independent controls and responses controls) are mapped into these NL2SOL internal controls. In particular, Dakota’s convergence_tolerance, max_iterations, max_function_evaluations, and fd_gradient_step_size controls are mapped directly into NL2SOL’s rfctol, mxiter, mxfcal, and dltfdj controls, respectively. In addition, Dakota’s fd_hessian_step_size is mapped into both delta0 and dltfdc, and Dakota’s output verbosity is mapped into NL2SOL’s auxxprt and outlev (for normal/verbose/debug output, NL2SOL prints initial guess, final solution, solution statistics, non-default values, and changes to the active bound constraint set on every iteration; for quiet output, NL2SOL prints only the initial guess and final solution; and for silent output, NL2SOL output is suppressed).
See Also

These keywords may also be of interest:

- `nlssol_sqp`
- `optpp_g_newton`
- `field_calibration_terms`

7.2.41.1 `function_precision`

- `Keywords Area`
- `method`
- `nl2sol`
- `function_precision`

Specify the maximum precision of the analysis code responses

**Specification**

*Alias*: none

*Argument(s)*: REAL

*Default*: 1.0e-10

**Description**

The `function_precision` control provides the algorithm with an estimate of the accuracy to which the problem functions can be computed. This is used to prevent the algorithm from trying to distinguish between function values that differ by less than the inherent error in the calculation.

7.2.41.2 `absolute_conv_tol`

- `Keywords Area`
- `method`
- `nl2sol`
- `absolute_conv_tol`

Absolute convergence tolerance

**Specification**

*Alias*: none

*Argument(s)*: REAL

*Default*: -1. (use NL2SOL internal default)
**7.2 method**

**Description**

\texttt{absolute\_conv\_tol} (NL2SOL's \texttt{afctol}) is the absolute function convergence tolerance (stop when half the sum of squares is less than \texttt{absolute\_conv\_tol}, which is mainly of interest on zero-residual test problems). The internal default is a function of machine epsilon (as limited by \texttt{function\_precision}). The default is selected with a value of -1.

**7.2.41.3 x\_conv\_tol**

- **Keywords Area**
- **method**
- **nl2sol**
- **x\_conv\_tol**

X-convergence tolerance

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** -1. (use NL2SOL internal default)

**Description**

\texttt{x\_conv\_tol} maps to the internal NL2SOL control \texttt{xctol}. It is the X-convergence tolerance (scaled relative accuracy of the solution variables).

The internal default is a function of machine epsilon (as limited by \texttt{function\_precision}). The default is selected with a value of -1.

**7.2.41.4 singular\_conv\_tol**

- **Keywords Area**
- **method**
- **nl2sol**
- **singular\_conv\_tol**

Singular convergence tolerance

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** -1. (use NL2SOL internal default)
singular_conv_tol (NL2SOL’s sctol) is the singular convergence tolerance, which works in conjunction with singular_radius to test for underdetermined least-squares problems (stop when the relative reduction yet possible in the sum of squares appears less than singular_conv_tol for steps of scaled length at most singular_radius).

The internal default is a function of machine epsilon (as limited by function_precision). The default is selected with a value of -1.

7.2.41.5 singular_radius

- Keywords Area
- method
- nl2sol
- singular_radius

Singular radius

Specification

Alias: none

Argument(s): REAL

Default: -1. (use NL2SOL internal default of 1)

Description

singular_radius works in conjunction with singular_conv_tol to test for underdetermined least-squares problems (stop when the relative reduction yet possible in the sum of squares appears less than singular_conv_tol for steps of scaled length at most singular_radius).

The internal default results in the internal use of steps of length 1. The default is selected with a value of -1.

7.2.41.6 false_conv_tol

- Keywords Area
- method
- nl2sol
- false_conv_tol

False convergence tolerance

Specification

Alias: none

Argument(s): REAL

Default: -1. (use NL2SOL internal default)
false_conv_tol (NL2SOL's xftol) is the false-convergence tolerance (stop with a suspicion of discontinuity when a more favorable stopping test is not satisfied and a step of scaled length at most false_conv_tol is not accepted)

The internal default is a function of machine epsilon (as limited by function_precision). The default is selected with a value of -1.

7.2.41.7 initial_trust_radius

- Keywords Area
- method
- nl2sol
- initial_trust_radius

Initial trust region radius

Specification

Alias: none
Argument(s): REAL
Default: -1. (use NL2SOL internal default of 1)

Description

initial_trust_radius specification (NL2SOL's lmax0) specifies the initial trust region radius for the algorithm.

The internal default results in the internal use of steps of length 1. The default is selected with a value of -1.

7.2.41.8 covariance

- Keywords Area
- method
- nl2sol
- covariance

Determine how the final covariance matrix is computed

Specification

Alias: none
Argument(s): INTEGER
Default: 0 (no covariance)
covariance (NL2SOL's covreq) specifies whether and how NL2SOL computes a final covariance matrix.

The desired covariance approximation:

- 0 = default = none
- 1 or -1 ==> \( \sigma^2 H^{-1} J^T J H^{-1} \)
- 2 or -2 ==> \( \sigma^2 H^{-1} \)
- 3 or -3 ==> \( \sigma^2 (J^T J)^{-1} \)
- Negative values ==> estimate the final Hessian \( H \) by finite differences of function values only (using \( \text{fd}_-\text{hessian_step_size} \))
- Positive values ==> differences of gradients (using \( \text{fd}_\text{hessian_step_size} \))

7.2.41.9 regression_diagnostics

- Keywords Area
- method
- nl2sol
- regression_diagnostics

Turn on regression diagnostics

Specification

Alias: none
Argument(s): none
Default: no regression diagnostics

Description

When regression_diagnostics (NL2SOL's rdreq) is specified and a positive-definite final Hessian approximation \( H \) is computed, NL2SOL computes and prints a regression diagnostic vector \( RD \) such that if omitting the \( i \)-th observation would cause \( \alpha \) times the change in the solution that omitting the \( j \)-th observation would cause, then \( RD[i] = |\alpha| RD[j] \). The finite-difference step-size tolerance affecting \( H \) is \( \text{fd}_\text{step_size} \) (NL2SOL's \( \text{delta}0 \) and \( \text{dltdfc} \)).

7.2.41.10 convergence_tolerance

- Keywords Area
- method
- nl2sol
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do. Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.41.11 max_iterations

- Keywords Area
- method
- nl2sol
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_{reliability, interval_est, evidence} / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max_function_evaluations

7.2.41.12 speculative

- Keywords Area
- method
- nl2sol
- speculative

Compute speculative gradients

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no speculation

**Description**

When performing gradient-based optimization in parallel, speculative gradients can be selected to address the load imbalance that can occur between gradient evaluation and line search phases. In a typical gradient-based
optimization, the line search phase consists primarily of evaluating the objective function and any constraints at a trial point, and then testing the trial point for a sufficient decrease in the objective function value and/or constraint violation. If a sufficient decrease is not observed, then one or more additional trial points may be attempted sequentially. However, if the trial point is accepted then the line search phase is complete and the gradient evaluation phase begins. By speculating that the gradient information associated with a given line search trial point will be used later, additional coarse grained parallelism can be introduced by computing the gradient information (either by finite difference or analytically) in parallel, at the same time as the line search phase trial-point function values. This balances the total amount of computation to be performed at each design point and allows for efficient utilization of multiple processors. While the total amount of work performed will generally increase (since some speculative gradients will not be used when a trial point is rejected in the line search phase), the run time will usually decrease (since gradient evaluations needed at the start of each new optimization cycle were already performed in parallel during the line search phase). Refer to [13] for additional details. The speculative specification is implemented for the gradient-based optimizers in the DOT, CONMIN, and OPT++ libraries, and it can be used with dakota numerical or analytic gradient selections in the responses specification (refer to responses gradient section for information on these specifications). It should not be selected with vendor numerical gradients since vendor internal finite difference algorithms have not been modified for this purpose. In full-Newton approaches, the Hessian is also computed speculatively. NPSOL and NLSSOL do not support speculative gradients, as their gradient-based line search in user-supplied gradient mode (dakota numerical or analytic gradients) is a superior approach for load-balanced parallel execution.

The speculative specification enables speculative computation of gradient and/or Hessian information, where applicable, for parallel optimization studies. By speculating that the derivative information at the current point will be used later, the complete data set (all available gradient/Hessian information) can be computed on every function evaluation. While some of these computations will be wasted, the positive effects are a consistent parallel load balance and usually shorter wall clock time. The speculative specification is applicable only when parallelism in the gradient calculations can be exploited by Dakota (it will be ignored for vendor numerical gradients).

7.2.41.13 max_function_evaluations

- **Keywords Area**
- **method**
- **nl2sol**
- **max_function_evaluations**

Number of function evaluations allowed for optimizers

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before `max_function_evaluations` was exceeded.

**Default Behavior**
Default value is 1000.

7.2.41.14  **scaling**
- **Keywords Area**
- **method**
- **nl2sol**
- **scaling**

Turn on scaling for variables, responses, and constraints

**Topics**
This keyword is related to the topics:
- **method_independent_controls**

**Specification**

**Alias:** none
**Argument(s):** none
**Default:** no scaling

**Description**
Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `*scales` keyword, which gives characteristic values (divisors)
- a `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**
There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval $[0,1]$;
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into $[0,1]$.

3. **log** - logarithmic scaling
   First, any characteristic values from the optional *_scales specification are applied. Then logarithm base 10 scaling is applied.
   Logarithmic scaling is not available for linear constraints.
   When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0e10*DBL_MIN$. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.41.15 **model_pointer**

- **Keywords Area**
- **method**
• nl2sol
• model_pointer

Identifier for model block to be used by a method

Topics
This keyword is related to the topics:

• block_pointer

Specification

**Alias:** none  
**Argument(s):** STRING  
**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.  
See `block_pointer` for details about pointers.

Examples

environment
  tabular_data  
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                         0.1 0.2 0.6
                         0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
7.2 method

model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.42 nonlinear_cg

  • Keywords Area

  • method

  • nonlinear_cg

(Experimental) nonlinear conjugate gradient optimization

Topics

This keyword is related to the topics:

  • local_optimization_methods

Specification

Alias: none

Argument(s): none

Child Keywords:
### Dakota Keyword Description

**Required/Optional Group**

<table>
<thead>
<tr>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>misc_options</strong></td>
<td>Options for nonlinear CG optimizer</td>
</tr>
<tr>
<td><strong>convergence_tolerance</strong></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td><strong>max_iterations</strong></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td><strong>scaling</strong></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td><strong>model_pointer</strong></td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

This method is an incomplete experimental implementation of nonlinear conjugate gradient optimization, a local, gradient-based solver.

**7.2.42.1 misc_options**

- **Keywords Area**
- **method**
- **nonlinear_cg**
- **misc_options**

Options for nonlinear CG optimizer

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** no misc options

**Description**

List of miscellaneous string options to pass to the experimental nonlinear CG solver (see NonlinearCGOptimizer.cpp in the Dakota source code for available controls). Includes controls for step sizes, linesearch control, convergence, etc.
7.2.42.2  convergence_tolerance

- Keywords Area
- method
- nonlinear_cg
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.42.3  max_iterations

- Keywords Area
- method
Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: $25\times n$)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \( \text{max\_iterations} \) iterations. In cases where there are multiple nested iterations to be controlled, \( \text{max\_iterations} \) generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- \( \text{max\_function\_evaluations} \)

7.2.42.4 scaling

- Keywords Area
- method
- nonlinear_cg
- scaling

Turn on scaling for variables, responses, and constraints

**Topics**

This keyword is related to the topics:

- method_independent_controls
**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no scaling

**Description**

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `∗scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `∗scales` keyword, which gives characteristic values (divisors)
- a `∗scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   
   First, any characteristic values from the optional `∗_scales` specification are applied. Then logarithm base 10 scaling is applied.

   Logarithmic scaling is not available for linear constraints.

   When continuous design variables are log scaled, linear constraints are not allowed.
Scales

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- *auto*, *log* - optional
- *value* - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN*. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

Examples

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

7.2.42.5 model_pointer

- Keywords Area
- method
- nonlinear_cg
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                  0.1 0.2 0.6
                  0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```
7.2.43 ncsu_direct

- Keywords Area
  - method
  - ncsu_direct

Dividing RECTangles method

Topics

This keyword is related to the topics:

- global_optimization_methods

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>solution_target</td>
<td></td>
<td>Specifies a globally optimal value toward which the optimizer should track</td>
</tr>
<tr>
<td>Optional</td>
<td>min_boxsize_limit</td>
<td></td>
<td>Stopping Criterion based on shortest edge of hyperrectangle</td>
</tr>
<tr>
<td>Optional</td>
<td>volume_boxsize_limit</td>
<td></td>
<td>Stopping criterion based on volume of search space</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
</tbody>
</table>
### Description

North Carolina State University (NCSU) has an implementation of the DIRECT algorithm (Dividing RECTangles algorithm that is outlined in the SCOLIB method section above). This version is documented in [28] We have found that the NCSU DIRECT implementation works better and is more robust for some problems than `coliny-direct`. Currently, we maintain both versions of DIRECT in Dakota; in the future, we may deprecate one.

The NCSU DIRECT method is selected with `ncsu_direct`. We have tried to maintain consistency between the keywords in SCOLIB and NCSU implementation of DIRECT, but the algorithms have different parameters, so the keywords sometimes have slightly different meaning.

### Stopping Criteria

The algorithm stops based on:

1. `max_iterations` - number of iterations
2. `max_function_evaluations` - number of function evaluations
3. `solution_target` and `convergence_tolerance`
4. `min_boxsize_limit`
5. `volume_boxsize_limit`

This method will always strictly respect the number of iterations, but may slightly exceed the number of function evaluations, as it will always explore all sub-rectangles at the current level.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions` are specified)
- Best Nonlinear Constraints
- Calibration (when `calibration_terms` are specified)

### See Also

These keywords may also be of interest:

- `coliny_direct`
7.2.43.1 solution_target

- Keywords Area
- method
- ncsu_direct
- solution_target

Specifies a globally optimal value toward which the optimizer should track.

**Specification**

**Alias:** solution_accuracy

**Argument(s):** REAL

**Default:** 0

**Description**

The solution target specifies a goal toward which the optimizer should track. This is used for test problems, when the true global minimum is known (call it solution_target := fglobal). Then, the optimization terminates when \(100(f_{\text{min}} - f_{\text{global}})/\max(1,\text{abs}(f_{\text{global}})) < \text{convergence_tolerance} \). The default for fglobal is -1.0e100 and the default for convergence tolerance is described at convergence_tolerance.

7.2.43.2 min_boxsize_limit

- Keywords Area
- method
- ncsu_direct
- min_boxsize_limit

Stopping Criterion based on shortest edge of hyperrectangle

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0e-4

**Description**

min_boxsize_limit is a setting that terminates the optimization when the measure of a hyperrectangle S with \(f(c(S)) = f_{\text{min}}\) is less than min_boxsize_limit. Each subregion considered by DIRECT has a size, which corresponds to the longest diagonal of the subregion. min_boxsize_limit specification terminates DIRECT if the size of the smallest subregion falls below this threshold. In practice, this specification is likely to be more effective at limiting DIRECT's search.
7.2.43.3 volume_boxsize_limit

- Keywords Area
- method
- ncsu_direct
- volume_boxsize_limit

Stopping criterion based on volume of search space

Specification

Alias: none
Argument(s): REAL
Default: 1.0e-6

Description

volume_boxsize_limit is a setting that terminates the optimization when the volume of a hyperrectangle S with f(c(S)) = \text{fmin} is less than volume_boxsize_limit percent of the original hyperrectangle. Basically, volume_boxsize_limit stops the optimization when the volume of the particular rectangle which has \text{fmin} is less than a certain percentage of the whole volume.

7.2.43.4 convergence_tolerance

- Keywords Area
- method
- ncsu_direct
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4
Description

The **convergence_tolerance** specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a **relative convergence tolerance** for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by **convergence_tolerance**, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- **DOT**: relative tolerance that must be satisfied for two consecutive iterations
- **NL2SOL**: See *nl2sol*
- **NLPQLP**: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- **NPSOL**: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.43.5 max_iterations

- **Keywords Area**
  - **method**
  - **ncsu_direct**
  - **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- **method_independent_controls**

Specification

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 100 (exceptions: fsu_cvt , local_reliability: 25; global_{reliability , interval_est , evidence} / efficient_global: 25*n)
7.2 method

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max function evaluations}

7.2.43.6 max_function_evaluations

- Keywords Area
- method
- ncsu_direct
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- \texttt{method independent controls}

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: INTEGER

\textbf{Default}: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max function evaluations} evaluations. See also \texttt{max iterations}.

Some optimizers (e.g., ncsu_direct) may run past this limit in the course of an iteration step that began before \texttt{max function evaluations} was exceeded.

Default Behavior

Default value is 1000.
7.2.43.7 scaling

- Keywords Area
- method
- ncsu_direct
- scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): none
Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling

   The target quantity will be scaled (divided) by the specified characteristic value.
2. **auto** - automatic scaling

First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:

- Two-sided bounds scaled into the interval \([0,1]\);
- One-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
- No bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into \([0,1]\).

3. **log** - logarithmic scaling

First, any characteristic values from the optional *scales specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.
When continuous design variables are log scaled, linear constraints are not allowed.

### Scales

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than \(1.0\times 10^{10}\times \text{DBL_MIN}\). User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

### Examples

The two examples below are equivalent:

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100
```

```
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

### 7.2.43.8 model_pointer

- **Keywords Area**
- **method**
- **ncsu_direct**
- **model_pointer**

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
    method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'
variables
uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = ’x1’ ’x2’

interface
  id_interface = ’I1’
  system async evaluation_concurrency = 5
              analysis_driver = ’text_book’

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.44 genie_opt_darts
  • Keywords Area
  • method
  • genie_opt_darts

Voronoi-based high-dimensional global Lipschitzian optimization

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td></td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>scaling</td>
<td></td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td></td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

Description

OPT-Darts method is a fast alternative to DIRECT for global Lipschitzian optimization purposes. Instead of hyperrectangular, OPT-Darts decomposes a high-dimensional domain into Voronoi cells, and places samples via stochastic blue noise instead of deterministic cell division.

To refine a cell, OPT-Darts first adds a new sample within it via spoke-dart sampling, then set the conflict radius to the cell’s inscribed hypersphere radius, to avoid adding a sample point that is too close to a prior sample, then divide
that cell (and update its neighboring cells) via the approximate Delaunay graph, and use the computed witnesses to
decide the next refinement candidate. These two steps replace the corresponding deterministic center-sample and
rectangular cell division in DIRECT, respectively.

OPT-Darts is the first exact stochastic Lipschitzian optimization technique that combines the benefits of guaranteed
convergence in [Jones et al. 1993] and high dimensional efficiency in [Spall 2005]. Computing blue noise and
Voronoi regions has been intractable in high dimensions, and are being done within OPT-Darts using Spoke-Darts.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to
HDF5:

- Best Parameters
- Best Objective Functions (when `objective_functions` are specified)
- Calibration (when `calibration_terms` are specified)

### 7.2.44.1 seed

- Keywords Area
- method
- genie_opt_darts
- seed

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of
the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value.
Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same
seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That
value can then be added to the Dakota input file.
Examples

```plaintext
method
   sampling
      sample_type lhs
      samples = 10
      seed = 15347
```

7.2.44.2 max_function_evaluations

- Keywords Area
- method
- genie_opt_darts
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max_function_evaluations} evaluations. See also \texttt{max_iterations}.

Some optimizers (e.g. \texttt{ncsu_direct}) may run past this limit in the course of an iteration step that began before \texttt{max_function_evaluations} was exceeded.

**Default Behavior**

Default value is 1000.

7.2.44.3 scaling

- Keywords Area
- method
- genie_opt_darts
- scaling

Turn on scaling for variables, responses, and constraints
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the `scaling` keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `∗scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a `∗scales` keyword, which gives characteristic values (divisors)
- a `∗scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:

1. value - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. auto - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Futher, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].
3. *log* - logarithmic scaling

First, any characteristic values from the optional *scales* specification are applied. Then logarithm base 10 scaling is applied.

Logarithmic scaling is not available for linear constraints.

When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales* keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types*, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- *auto, log* - optional
- *value* - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than $1.0\times 10^{-10}$ DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = "value"
   primary_scales = 1 1 100

responses
   objective_functions 3
   sense "maximize"
   primary_scale_types = "value" "value" "value"
   primary_scales = 1 1 100
```

7.2.44.4 model_pointer

- **Keywords Area**
- **method**
- **genie_opt_darts**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
  samples = 10
  seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
  lower_bound = 0. 0.
  upper_bound = 1. 1.
  descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
```
7.2 method

```
analysis_driver = 'text_book'
responses
response_functions = 3
no_gradients
no_hessians
```

7.2.45 genie_direct

- Keywords Area
- method
- genie_direct

Classical high-dimensional global Lipschitzian optimization

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_function_evaluations</td>
<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scaling</td>
<td>Turn on scaling for variables, responses, and constraints</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

DIRECT (Dlviding RECTangles) partitions the domain into hyperrectangles and uses an iterative Lipschitzian optimization approach to search for a global optimal point.

DIRECT begins by scaling the domain into the unit hypercube by adopting a center-sampling strategy. The objective function is evaluated at the midpoint of the domain, where a lower bound is constructed. In one-dimension, the domain is tri-sected and two new center points are sampled. At each iteration (dividing and sampling), DIRECT identifies intervals that contain the best minimal value of the objective function found up to that point. This strategy of selecting and dividing gives DIRECT its performance and convergence properties compared to other deterministic methods.

The classical DIRECT method [Shubert 1972] has two limitations: poor scaling to high dimensions; and relying on a global K; whose exact value is often unknown. The enhanced DIRECT algorithm [Jones et al. 1993] generalizes [Shubert 1972] to higher dimensions and does not require knowledge of the Lipschitz constant.
7.2.45.1 seed

- Keywords Area
- method
- genie_direct
- seed

Seed of the random number generator

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.45.2 max_function_evaluations

- Keywords Area
- method
- genie_direct
- max_function_evaluations

Number of function evaluations allowed for optimizers
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed \texttt{max\_function\_evaluations} evaluations. See also \texttt{max\_iterations}.

Some optimizers (e.g. \texttt{ncsu\_direct}) may run past this limit in the course of an iteration step that began before \texttt{max\_function\_evaluations} was exceeded.

Default Behavior

Default value is 1000.

7.2.45.3 scaling

Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the \texttt{scaling} keyword. Discrete variable scaling is not supported.
When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords `scale_types` and/or `*scales` in the `variables` and `responses` blocks. When the `scaling` keyword is omitted from `method`, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- A `*scales` keyword, which gives characteristic values (divisors)
- A `*scale_types` keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in `responses` for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

**Scale Types**

There are three scale types:

1. **value** - characteristic value scaling
   - The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   - First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
     - two-sided bounds scaled into the interval [0,1];
     - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
     - no bounds or targets: no automatic scaling possible, therefore no scaling for this component
   - Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   - First, any characteristic values from the optional `*_scales` specification are applied. Then logarithm base 10 scaling is applied.
   - Logarithmic scaling is not available for linear constraints.
   - When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The `*scales` keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding `*scale_types`, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- **auto, log** - optional
- **value** - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than `1.0e10*DBL_MIN`. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.
Examples

The two examples below are equivalent:

```plaintext
responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value"
    primary_scales = 1 1 100

responses
    objective_functions 3
    sense "maximize"
    primary_scale_types = "value" "value" "value"
    primary_scales = 1 1 100
```

7.2.45.4 model_pointer

- Keywords Area
- method
- genie_direct
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.46 efficient_global

- Keywords Area
- method
- efficient_global

Global Surrogate Based Optimization, a.k.a. EGO

Topics

This keyword is related to the topics:

- global_optimization_methods
- surrogate_based_optimization_methods
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<th>Dakota Keyword</th>
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<td></td>
<td>initial_samples</td>
<td>Initial number of samples for sampling-based methods</td>
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<td>seed</td>
<td>Seed of the random number generator</td>
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<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
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<td>Expected improvement convergence tolerance</td>
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<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
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<td>File containing points you wish to use to build a surrogate</td>
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<td>export_approx_-points_file</td>
<td>Output file for surrogate model value evaluations</td>
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</table>
Description

The Efficient Global Optimization (EGO) method was first developed by Jones, Schonlau, and Welch[55]. In EGO, a stochastic response surface approximation for the objective function is developed based on some sample points from the “true” simulation.

Note that several major differences exist between our implementation and that of[55]. First, rather than using a branch and bound method to find the point which maximizes the EIF, we use the DIRECT global optimization method.

Second, we support both global optimization and global nonlinear least squares as well as general nonlinear constraints through abstraction and subproblem recasting.

The efficient global method is in prototype form. Currently, we do not expose any specification controls for the underlying Gaussian process model used or for the optimization of the expected improvement function (which is currently performed by the NCSU DIRECT algorithm using its internal defaults).

By default, EGO uses the Surfpack GP (Kriging) model, but the Dakota implementation may be selected instead. If use_derivatives is specified the GP model will be built using available derivative data (Surfpack GP only).

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Best Parameters
- Best Objective Functions (when objective_functions) are specified)
- Best Nonlinear Constraints
- Calibration (when calibration_terms are specified)

Theory

The particular response surface used is a Gaussian process (GP). The GP allows one to calculate the prediction at a new input location as well as the uncertainty associated with that prediction. The key idea in EGO is to maximize the Expected Improvement Function (EIF). The EIF is used to select the location at which a new training point should be added to the Gaussian process model by maximizing the amount of improvement in the objective function that can be expected by adding that point. A point could be expected to produce an improvement in the objective function if its predicted value is better than the current best solution, or if the uncertainty in its prediction is such that the probability of it producing a better solution is high. Because the uncertainty is higher in regions of the design space with few observations, this provides a balance between exploiting areas of the design space that predict good solutions, and exploring areas where more information is needed. EGO trades off this “exploitation vs. exploration.”

The general procedure for these EGO-type methods is:

- Build an initial Gaussian process model of the objective function
- Find the point that maximizes the EIF. If the EIF value at this point is sufficiently small, stop.
- Evaluate the objective function at the point where the EIF is maximized. Update the Gaussian process model using this new point.
- Return to the previous step.
See Also

These keywords may also be of interest:

- surrogate_based_local
- surrogate_based_global

7.2.46.1 initial_samples

- Keywords Area
- method
- efficient_global
- initial_samples

Initial number of samples for sampling-based methods

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** \((d+1)(d+2)/2\)

Description

The \texttt{initial_samples} keyword is used to define the number of initial samples (i.e., randomly chosen sets of variable values) at which to execute a model. The initial samples may later be augmented in an iterative process.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least \(d+1\) samples should be used, where \(d\) is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \((d+1)(d+2)/2\) samples are needed. For uncertainty quantification, we recommend at least \(10d\) samples. For \texttt{variance_based_decomp}, we recommend hundreds to thousands of samples. Note that for \texttt{variance_based_decomp}, the number of simulations performed will be \(N=(d+2)\).

Examples

```plaintext
method
ing sampling
  sample_type random
  initial_samples = 20
  refinement_samples = 5

7.2.46.2 seed

- Keywords Area
- method
- efficient_global
```
• **seed**

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

### 7.2.46.3 batch_size

- **Keywords Area**
- **method**
- **efficient_global**
- **batch_size**

Total batch size in parallel EGO

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Child Keywords:**
### Description

Refinement candidates are generated by an acquisition function such as maximum expected improvement, which balances exploration and exploitation. Refinement candidates can also be generated by purely explorative metrics such as maximum prediction variance. For a specified `batch_size`, `exploration` specifies the subset of this total that will be dedicated to pure exploration of the parameter space.

**Default Behavior** All of the batch size is devoted to the standard acquisition approach, balancing exploration and exploitation.

### Examples

```plaintext
method,  
    efficient_global  
    seed = 1237
batch_size = 8  # total
exploration = 2  # 2 out of 8
```

### 7.2.46.3.1 exploration

- **Keywords Area**
  - method
  - `efficient_global`
  - `batch_size`
  - `exploration`

Portion of batch size dedicated to exploration in parallel EGO

### Specification

**Alias:** none

**Argument(s):** INTEGER

### Description

Refinement candidates are generated by an acquisition function such as maximum expected improvement, which balances exploration and exploitation. Refinement candidates can also be generated by purely explorative metrics such as maximum prediction variance.
such as maximum prediction variance. For a specified batch_size, exploration specifies the subset of this total that will be dedicated to pure exploration of the parameter space.

**Default Behavior** All of the batch size is devoted to the standard acquisition approach, balancing exploration and exploitation.

### Examples

```python
method,
  efficient_global
  seed = 1237
  batch_size = 8  # total
  exploration = 2  # 2 out of 8
```

#### 7.2.46.3.2 synchronization

- **Keywords Area**
- **method**
- **efficient_global**
- **batch_size**
- **synchronization**

Select how Dakota schedules a batch of concurrent function evaluations in a parallel algorithm

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** method specific

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Description</th>
</tr>
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<td></td>
<td></td>
<td>blocking</td>
<td>Complete concurrent evaluation of all evaluations within a batch</td>
</tr>
<tr>
<td></td>
<td></td>
<td>nonblocking</td>
<td>A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm</td>
</tr>
</tbody>
</table>

**Description**

The synchronization specification can be used to specify the use of either blocking or nonblocking schedulers, where the former must block until all evaluations are completed and the latter does not block, returning only the partial set that has been completed for a particular call to the scheduler.
7.2.46.3.2.1 blocking

- Keywords Area
- method
- efficient_global
- batch_size
- synchronization
- blocking

Complete concurrent evaluation of all evaluations within a batch

Specification

**Alias:** none

**Argument(s):** none

Description

In the **blocking** case, all of the concurrent evaluations in the batch must be completed and all results are returned together to the algorithm.

These runs are reproducible, assuming use of the same seed in cases with random number generation.

7.2.46.3.2.2 nonblocking

- Keywords Area
- method
- efficient_global
- batch_size
- synchronization
- nonblocking

A batch of concurrent evaluations are queried for completions and a partial set is returned to the algorithm

Specification

**Alias:** none

**Argument(s):** none

Description

In the **nonblocking** case, the set of running evaluations is queried for completion and a partial set of results is returned to an asynchronous algorithm, which can then generate new evaluations based on these partial results.

In all cases, the algorithm steps will be subject to parallel timing variabilities / race conditions such that these runs will not generally be repeatable regardless of seed specification.
7.2.46.4 max_iterations

• Keywords Area
• method
• efficient_global
• max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations

7.2.46.5 convergence_tolerance

• Keywords Area
• method
• efficient_global
• convergence_tolerance

Expected improvement convergence tolerance
7.2 method

Specification

Alias: none
Argument(s): REAL
Default: 1.e-12

Description

The Efficient Global Optimization method will terminate if the expected improvement is below the specified tolerance for two subsequent iterations.

7.2.46.6  x_conv_tol

- Keywords Area
- method
- efficient_global
- x_conv_tol

x-convergence tolerance

Specification

Alias: none
Argument(s): REAL
Default: 1.0e-8

Description

The Efficient Global Optimization method will terminate if the relative change in the best decision variables values x is less than the specified tolerance.

7.2.46.7  gaussian_process

- Keywords Area
- method
- efficient_global
- gaussian_process

Gaussian Process surrogate model

Specification

Alias: kriging
Argument(s): none
Default: Surfpack Gaussian process
Child Keywords:
Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

### 7.2.46.7.1 surfpack

- Keywords Area
  - method
  - efficient_global
  - gaussian_process
  - surfpack

Use the Surfpack version of Gaussian Process surrogates

### Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
7.2 method

Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See optimization_method.

   The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See trend.

3. Correlation Lengths:
   
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See correlation_lengths.

4. Ill-conditioning:

   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.

   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.

   The surfpack model handles ill-conditioning internally by default, but behavior can be modified using

5. Gradient Enhanced Kriging (GEK):

   The use_derivatives keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).

   See notes in the Theory section.

Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.
7.2.46.7.1.1 export_model

• Keywords Area
• method
• efficient_global
• gaussian_process
• surfpack
• export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
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</tr>
<tr>
<td>Required</td>
<td></td>
<td>formats</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

• text_archive - Plain-text, machine-readable archive for re-import or use with the surfpack executable
• binary_archive - Binary, machine-readable archive for re-import or use with the surfpack executable
• algebraic_file - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
• algebraic_console - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable
These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

**Default Behavior**

No export.

**Expected Output**

Output depends on selected format; see the formats specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.46.7.1.2 filename_prefix

- Keywords Area
- method
- efficient_global
- gaussian_process
- surfpack
- export_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

- `surrogate_models`

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```
surrogate global gaussian_process surfpack
   export_model
      filename_prefix = 'my_surrogate'
      formats
         algebraic_file
         binary_archive
```

7.2.46.7.1.3 formats

- Keywords Area
- method
- efficient_global
- gaussian_process
- surfpack
- export_model
- formats

Formats for surrogate model export

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Dakota Keyword Description</th>
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</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See `export_model` and the entries for the format selection keywords for further information.

#### 7.2.46.7.1.4 text_archive

- **Keywords Area**
- **method**
- **efficient_global**
- **gaussian_process**
- **surfpack**
- **export_model**
- **formats**
- **text_archive**

Surrogate model plain-text archive file format

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.
Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.46.7.1.5 binary_archive

- Keywords Area
- method
- efficient_global
- gaussian_process
- surfpack
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin. Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which ‘bsps’ stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
• dakota

Select the built in Gaussian Process surrogate

Specification

Alias: none

Argument(s): none

Description

A second version of GP surrogates was available in prior versions of Dakota. For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in [60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

7.2.46.7.3 experimental

• Keywords Area
• method
• efficient_global
• gaussian_process
• experimental

Use the experimental Gaussian Process surrogate

Specification

Alias: none

Argument(s): none

Child Keywords:
The experimental GP surrogate is an emerging capability intended to supersede the *dakota* and *surfpack* GP implementations by unifying the best features of both. It also includes additional features such as the ability to export the model and then use it from Python.

**See Also**

These keywords may also be of interest:

- `experimental_gaussian_process`

**7.2.46.7.3.1 export_model**

- `Keywords Area`
- `method`
- `efficient_global`
- `gaussian_process`
- `experimental`
- `export_model`

Exports surrogate model in user-specified format(s)

**Topics**

This keyword is related to the topics:

- `surrogate_models`

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:
Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the `formats` specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive**: Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive**: Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file**: Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console**: Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Funtions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior

No export.

Expected Output

Output depends on selected format; see the `formats` specification.

Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>User-customizable portion of exported/imported surrogate model filenames</td>
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<tr>
<td>Required</td>
<td></td>
<td>formats</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>
• `gaussian_process`
• `experimental`
• `export_model`
• `filename_prefix`

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

• `surrogate_models`

**Specification**

**Alias**: none

**Argument(s)**: STRING

**Default**: exported_surrogate

**Description**

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

**Examples**

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```dakota
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive
```

---

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Formats for surrogate model export

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<td>Optional</td>
<td>binary_archive</td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See export_model and the entries for the format selection keywords for further information.

7.2.46.7.3.4 text_archive

- Keywords Area
  - method
  - efficient_global
  - gaussian_process
  - experimental
  - export_model
  - formats
  - text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models
Specification

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.response_descriptor.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.response_descriptor.sps`, in which `sps` stands for Surfpack surrogate. See `filename_prefix` for further information about surrogate file naming.

**Examples**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.46.7.3.5 `binary_archive`

- **Keywords Area**
- **method**
- **efficient_global**
- **gaussian_process**
- **experimental**
- **export_model**
- **formats**
- **binary_archive**

Surrogate model binary archive file format

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named `{prefix}.response_descriptor.bin`. 

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Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.46.7.3.6 options_file

- Keywords Area
- method
- efficient_global
- gaussian_process
- experimental
- options_file

Filename for a YAML file that specifies Gaussian process options

Specification

Alias: none

Argument(s): STRING

Default: no advanced options file

Description

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

Examples

```yaml
# FILE: gp_options.yaml
GP Parameters:
Sigma Bounds:
  lower bound: 1.0e-2
  upper bound: 1.0e2
scaler name: standardization
num restarts: 20
Nugget:
  fixed nugget: 0.0
  estimate nugget: true
Bounds:
  lower bound: 3.17e-8
  upper bound: 1.0e-2
gp seed: 42
standardize response: false
Trend:
  estimate trend: true
Options:
  max degree: 2
  reduced basis: false
```
p-norm: 1.0
scaler type: none
regression solver type: SVD
verbosity: 1
kernel type: squared exponential
Length-scale Bounds:
lower bound: 1.0e-2
upper bound: 1.0e2
verbosity: 1

7.2.46.8 use_derivatives

- Keywords Area
- method
- efficient_global
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.46.9 import_build_points_file

- Keywords Area
- method
- efficient_global
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file
Argument(s): STRING
Default: no point import from a file
Child Keywords:
The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- `annotated` (default)
- `custom_annotated`
- `freeform`

**Examples**

```dakota
method
polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.46.9.1  `custom_annotated`

- **Keywords Area**
-  `method`
-  `efficient_global`
-  `import_build_points_file`
-  `custom_annotated`

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom.annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
7.2 method

```python
evironment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.46.9.1.1 header

- Keywords Area
- method
- efficient_global
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

7.2.46.9.1.2 eval_id

- Keywords Area
- method
- efficient_global
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column
Description

See description of parent custom_annotated

7.2.46.9.1.3 interface_id

• Keywords Area
• method
• efficient_global
• import_build_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.46.9.2 annotated

• Keywords Area
• method
• efficient_global
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.46.9.3 freeform

- Keywords Area
- method
- efficient_global
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
```

7.2.46.9.4 active_only

- **Keywords Area**

- **method**

- **efficient_global**

- **import_build_points_file**

- **active_only**

Import only active variables from tabular data file
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables. This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.46.10 export_approx_points_file

- Keywords Area
- method
- efficient_global
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
### Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

### Usage Tips

Dakota exports tabular data in one of three formats:

- `annotated` (default)
- `custom_annotated`
- `freeform`

#### 7.2.46.10.1 custom_annotated

- **Keywords Area**
- **method**
- **efficient_global**
- `export_approx_points_file`
- `custom_annotated`

Selects custom-annotated tabular file format

### Topics

This keyword is related to the topics:

- `file_formats`

### Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td><code>header</code></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.46.10.1.1 header

- Keywords Area
- method
- efficient_global
- export_approx_points_file
**custom_annotated**

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent `custom_annotated`.

**7.2.46.10.1.2 eval_id**

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`.

**7.2.46.10.1.3 interface_id**

Enable interface ID column in custom-annotated tabular file

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7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Default**: no interface_id column

Description

See description of parent custom.annotated

7.2.46.10.2 annotated

- Keywords Area
- method
- efficient_global
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias**: none

**Argument(s)**: none

**Default**: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom.annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated format`, though `freeform` remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id      interface    x1      x2     obj_fn  nln_ineq_con_1  nln_ineq_con_2
1             NO_ID       0.9     1.1    0.0002  0.26             0.76
2             NO_ID       0.90009 1.1    0.0001996404857 0.2601620081 0.759955
3             NO_ID       0.89991 1.1    0.0002003604863 0.2598380081 0.760045
...
```

7.2.46.10.3 freeform

- Keywords Area
- method
- efficient_global
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior
The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to **annotated** format, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In **freeform**, the \( \text{num\_rows} \times \text{num\_cols} \) total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.46.11 **model_pointer**

- **Keywords Area**

- **method**

- **efficient_global**

- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians
```
7.2.47 surrogate_based_uq

- Keywords Area
- method
- surrogate_based_uq

Generic UQ method for constructing and interrogating a surrogate model.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>samples_on_emulator</td>
<td>samples_on_emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>rng</td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>probability_refinement</td>
<td>probability_refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
<tr>
<td>--------------------------</td>
<td>----------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
</tr>
<tr>
<td>Optional(Choose One)</td>
<td>Covariance Type (Group 1)</td>
<td>diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td>Optional</td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

As surrogate models by stochastic expansion migrate into the model specification, this method provides a general-purpose UQ method to interrogate the surrogate for generating statistics. This method must identify the surrogate of interest through its `model_pointer`, distinguishing it from fully-
integrated method specifications such as polynomial_chaos, stoch_collocation, and function_train that couple directly with a simulation model (and form the PCE, SC, FT surrogate approximations implicitly prior to using them for generating statistics).

See Also

These keywords may also be of interest:

- polynomial_chaos
- stoch_collocation
- function_train

7.2.47.1  samples_on_emulator

- Keywords Area
- method
- surrogate_based_uq
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples

Argument(s): INTEGER

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
    polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.47.2  sample_type

- Keywords Area
- method
- surrogate_based_uq
- sample_type

Selection of sampling strategy
Specification

**Alias:** none

**Argument(s):** none

**Default:** random

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
</tbody>
</table>

**random**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

7.2.47.2.1 lhs

- Keywords Area
- method
- surrogate_based_uq
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias:** none

**Argument(s):** none
Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```plaintext
method sampling
  sample_type lhs
  samples = 20
```

**7.2.47.2.2 random**

- **Keywords Area**
  - `method`
  - `surrogate_based_uq`
  - `sample_type`
  - `random`

Uses purely random Monte Carlo sampling to sample variables.

**Specification**

- **Alias:** none
- **Argument(s):** none

**Description**

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.
Examples

method
  sampling
    sample_type random
    samples = 200

7.2.47.3  rng

  • Keywords Area
  • method
  • surrogate_based_uq
  • rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none
Default: Mersenne twister ( mt19937 )

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

Description

The rng keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the rng keyword must be accompanied by either rnum2 (pseudo-random numbers) or mt19937 (random numbers generated by the Mersenne twister). Otherwise, mt19937, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
7.2 method

7.2.47.3.1 mt19937

- Keywords Area
- method
- surrogate_based_uq
- rng
- mt19937

Generates random numbers using the Mersenne twister

Specification

Alias: none
Argument(s): none

Description

The mt19937 keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: http://en.wikipedia.org/wiki/Mersenne_twister.

Default Behavior

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.47.3.2 rnum2

- Keywords Area
- method
- surrogate_based_uq
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

Alias: none
Argument(s): none
Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.47.4 probability_refinement

- Keywords Area
- method
- surrogate_based_uq
- probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

- reliability_methods

Specification

**Alias:** sample_refinement

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Importance Sampling Approach (Group 1)</td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>
## 7.2 method

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>

### Optional

<table>
<thead>
<tr>
<th>Argument(s)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design.</td>
</tr>
</tbody>
</table>

### Description

The `probability_refinement` allows refinement of probability and generalized reliability results using importance sampling. If one specifies `probability_refinement`, there are some additional options. One can specify which type of importance sampling to use (`import`, `adapt_import`, or `mm_adapt_import`). Additionally, one can specify the number of refinement samples to use with `refinement_samples` and the seed to use with `seed`.

The `probability_refinement` density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

#### 7.2.47.4.1 import

- **Keywords Area**
- **method**
- **surrogate_based_uq**
- **probability_refinement**
- **import**

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

`import` centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

#### 7.2.47.4.2 adapt_import

- **Keywords Area**
- **method**
- **surrogate_based_uq**
- **probability_refinement**
- adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

7.2.47.4.3 mm_adapt_import

- **Keywords Area**  
  - method  
  - surrogate_based_uq  
  - probability_refinement  
  - mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).

7.2.47.4.4 refinement_samples

- **Keywords Area**  
  - method  
  - surrogate_based_uq  
  - probability_refinement  
  - refinement_samples

Number of samples used to refine a probability estimate or sampling design.
7.2 method

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial samples in a sampling design.

7.2.47.5 **final_moments**

- Keywords Area
- method
- surrogate_based_uq
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

**Default:** standard

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>
When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central

Overiding the default to none as follows:

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none
```n
allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```n
to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```n
7.2.47.5.1 none

- Keywords Area
- method
- surrogate_based_uq
- final_moments
• none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none

7.2.47.5.2 standard

• Keywords Area
• method
• surrogate_based_uq
• final_moments
• standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.
Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

7.2.47.6  response_levels

• **Keywords Area**

• method

• surrogate_based_uq

• response_levels

Values at which to estimate desired statistics for each response
7.2 method Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_response_</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>levels</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>compute</td>
<td>Selection of</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>statistics to</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>compute at each</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num-`
The Dakota input file below specifies a sampling method with response levels of interest.

```
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.60e+11 4.00e+11 4.40e+11
                   6.00e+04 6.30e+04 7.00e+04
                   3.50e+05 4.00e+05 4.50e+05

variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n', 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u', 'TF2u'
  weibull_uncertain = 2
  alphas = 12.1, 30.
  betas = 250.1, 590.
  descriptors = 'TF1w', 'TF2w'
  histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h', 'TF2h'

histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'

interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

```
Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
   Bin Lower  Bin Upper   Density Value
     ------  -------  ------------
 2.7604749078e+11 3.6000000000e+11  5.3601733194e-12
 3.6000000000e+11 4.0000000000e+11  4.2500000000e-12
 4.0000000000e+11 4.4000000000e+11  3.7500000000e-12
 4.4000000000e+11 5.4196114379e+11  2.2557612778e-12

PDF for response_fn_2:
   Bin Lower  Bin Upper   Density Value
     ------  -------  ------------
 4.6431154744e+04 6.0000000000e+04  2.8742313912e-05
 6.0000000000e+04 6.5000000000e+04  6.4000000000e-05
 6.5000000000e+04 7.0000000000e+04  4.0000000000e-05
 7.0000000000e+04 7.8702465755e+04  1.0341896485e-05

PDF for response_fn_3:
   Bin Lower  Bin Upper   Density Value
     ------  -------  ------------
 2.3796737090e+05 3.5000000000e+05  4.2844608686e-06
 3.5000000000e+05 4.0000000000e+05  8.6000000000e-06
```
Level mappings for each response function:

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.47.6.1 `num_response_levels`

- **Keywords Area**
  - method
  - surrogate_based_uq
  - response_levels
  - num_response_levels

Number of values at which to estimate desired statistics for each response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

**Description**

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.
**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```plaintext
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5

7.2.47.6.2 compute
  • Keywords Area
  • method
  • surrogate_based_uq
  • response_levels
  • compute
```

Selection of statistics to compute at each response level

**Specification**

*Alias*: none

*Argument(s)*: none

*Default*: probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>probabilities</td>
<td></td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
<td></td>
</tr>
<tr>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
<td></td>
</tr>
</tbody>
</table>
Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities, reliabilities, or gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

7.2.47.6.2.1 probabilities

- Keywords Area

- method

- surrogate_based_uq

- response_levels

- compute

- probabilities

Computes probabilities associated with response levels

Specification

**Alias**: none

**Argument(s)**: none
Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
    compute probabilities
```

7.2.47.6.2.2 `reliabilities`

- Keywords Area
- method
- surrogate_based_uq
- response_levels
- compute
- reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The `reliabilities` keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the reliabilities are not computed by default. To change this behavior, the `reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
7.2 method

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

7.2.47.6.2.3 gen_reliabilities

- Keywords Area
- method
- surrogate_based_uq
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```
Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

Description

With the system probability/reliability option, statistics for specified response levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.47.6.2.5 series

- Keywords Area
- method
- surrogate_based_uq
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system
Specification

**Alias:** none

**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.47.6.2.6 `parallel`

- `Keywords Area`
- `method`
- `surrogate_based_uq`
- `response_levels`
- `compute`
- `system`
- `parallel`

Aggregate response statistics assuming a parallel system

Specification

**Alias:** none

**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.47.7 `probability_levels`

- `Keywords Area`
- `method`
- `surrogate_based_uq`
- `probability_levels`

Specify probability levels at which to estimate the corresponding response value

Specification

**Alias:** none

**Argument(s):** `REALLIST`

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**
### Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

### Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

### Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method,
sampling,
samples = 100 seed = 1
complementary distribution
probability_levels = 1. .66 .33 0.
1. .8 .5 0.
1. .3 .2 0.
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
interface,
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'
responses,
response_functions = 3
```

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_-</td>
<td></td>
<td>Specify which probability_-levels correspond to which response</td>
</tr>
<tr>
<td></td>
<td>levels</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond to the probability levels specified in the input file. Those response values are also shown in the CCDF.

**Probability Density Function (PDF) histograms for each response function:**

**PDF for response_fn_1:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

**PDF for response_fn_2:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

**PDF for response_fn_3:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

**Level mappings for each response function:**

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

**7.2.47.7.1 num_probability_levels**

- **Keywords Area**
  - **method**
**Keywords Area**

- `surrogate_based_uq`
- `probability_levels`
- `num_probability_levels`

Specify which `probability_levels` correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** `probability_levels` evenly distributed among response functions

**Description**

See parent page

7.2.47.8 reliability_levels

- **Keywords Area**
- `method`
- `surrogate_based_uq`
- `reliability_levels`

Specify reliability levels at which the response values will be estimated

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><code>num_reliability_</code>- <code>levels</code></td>
<td>Specify which <code>reliability_</code>- <code>levels</code> correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.47.8.1 num_reliability_levels

- Keywords Area
- method
- surrogate_based_uq
- reliability_levels
- num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.47.9 gen_reliability_levels

- Keywords Area
- method
- surrogate_based_uq
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:
Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.47.9.1 num_gen_reliability_levels

- Keywords Area
- method
- surrogate_based_uq
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.47.10 distribution

- Keywords Area
- method
Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** cumulative (CDF)  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```plaintext
method
    sampling
        sample_type lhs
        samples = 10
        distribution cumulative
```

7.2.47.10.1 cumulative

- Keywords Area
- method
- surrogate_based_uq

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• distribution

• cumulative

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.47.10.2 complementary

• Keywords Area

• method

• surrogate_based_uq

• distribution

• complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary

7.2.47.11 variance_based_decomp

- Keywords Area
- method
- surrogate_based_uq
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects.

Specification

**Alias:** none

**Argument(s):** none

**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
</tbody>
</table>
Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

Default Behavior

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

Expected Outputs

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.

Examples

```plaintext
method,
  polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].
7.2 method

- variance_based_decomp
- interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

Description

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.47.11.2 drop_tolerance

- Keywords Area
- method
  - surrogate_based_uq
  - variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed

Description

The drop_tolerance keyword allows the user to specify a value below which sensitivity indices generated by variance_based_decomp are not displayed.

Default Behavior

By default, all sensitivity indices generated by variance_based_decomp are displayed.

Usage Tips

For polynomial_chaos, which outputs main, interaction, and total effects by default, the univariate_effects may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
Examples

method,
sampling
    sample_type lhs
    samples = 100
    variance_based_decomp
drop_tolerance = 0.001

7.2.47.12 diagonal_covariance

- Keywords Area
- method
- surrogate_based_uq
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none

Argument(s): none

Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.47.13 full_covariance

- Keywords Area
- method
- surrogate_based_uq
- full_covariance

Display the full covariance matrix

Specification

Alias: none

Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.
7.2.47.14  import_approx_points_file

- Keywords Area
- method
- surrogate_based_uq
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

**Specification**

**Alias:** none  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional</strong> (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

**Default Behavior**  No import of points at which to evaluate the surrogate.

**Expected Output**  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```
method
  polynomial_choas
    expansion_order = 4
    import_approx_points_file = 'import.mcmc_annot.dat'
      annotated
```

7.2.47.14.1  custom_annotated

- Keywords Area
- method
- surrogate_based_uq
Keywords Area

- `import_approx_points_file`
- `custom_annotated`

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- `file_formats`

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id

Resulting tabular file:
```

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.47.14.1.1 header

- Keywords Area
- method
- surrogate_based_uq
- import_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.47.14.1.2 eval_id

- Keywords Area
- method
- surrogate_based_uq
- import_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent `custom_annotated`

7.2.47.14.1.3 `interface_id`

- Keywords Area
- method
- surrogate_based_uq
- import_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent `custom_annotated`

7.2.47.14.2 `annotated`

- Keywords Area
- method
- surrogate_based_uq
- import_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
7.2 method

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

**Examples**

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

...
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9    1.1   0.0002    0.26    0.76
0.90009 1.1  0.0001996404857 0.2601620081 0.759955
0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...
```

7.2.47.14.4  active_only

- Keywords Area
• method
• surrogate_based_uq
• import_approx_points_file
• active_only

Import only active variables from tabular data file

Topics
This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description
By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.47.15 export_approx_points_file

• Keywords Area
• method
• surrogate_based_uq
• export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file
Argument(s): STRING
Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.47.15.1 custom_annotated

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2.47.15.1.1 header

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

7.2.47.15.1.2 eval_id

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent custom_annotated
7.2.47.15.1.3 interface_id

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent custom_annotated

7.2.47.15.2 annotated

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
    tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.47.15.3 freeform

- Keywords Area
- method
- surrogate_based_uq
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<p>| | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
<td></td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
<td></td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.47.16  seed

- **Keywords Area**
- **method**
- **surrogate_based_uq**
- **seed**

Seed of the random number generator
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.47.17  **fixed_seed**

- **Keywords Area**
- **method**
- **surrogate_based_uq**
- **fixed_seed**

Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run

Description

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The
same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the **seed** must also be specified.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

#### 7.2.47.18  **model_pointer**

- **Keywords Area**
- **method**
- **surrogate_based_uq**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

**Alias:** none  
**Argument(s):** STRING  
**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The **model_pointer** is used to specify which **model** block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a **model** block in the Dakota input file that has a corresponding **id_model** with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a **model_pointer** for each method is imperative.  
See **block_pointer** for details about pointers.
Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
  samples = 10
  seed = 98765 rng rnum2
  response_levels = 0.1 0.2 0.6
  0.1 0.2 0.6
  0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.48  function_train

  • Keywords Area
  • method
  • function_train

UQ method leveraging a functional tensor train surrogate model.

Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_- - iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_- - tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>metric_scale</td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>regression_type</td>
<td>Type of solver for forming function train approximations by regression</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_solver_- - iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cross_- - iterations</td>
<td>Maximum number of iterations for cross-approximation during a rank adaptation.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>solver_tolerance</td>
<td>Convergence tolerance for the optimizer used during the regression solve.</td>
</tr>
<tr>
<td>Optional</td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation</td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>------------------------</td>
<td>------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion. Number of collocation points used to estimate expansion coefficients</td>
<td></td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Collocation Control (Group 1)</td>
<td>collocation_points</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rounding_tolerance</td>
<td>An accuracy tolerance that is used to guide rounding during rank adaptation</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>arithmetic_tolerance</td>
<td>A secondary rounding tolerance used for post-processing</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>start_order</td>
<td>(Initial) polynomial order of each univariate function within the functional tensor train</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>adapt_order</td>
<td>Activate adaptive procedure for determining the best basis order</td>
<td></td>
</tr>
<tr>
<td>Field</td>
<td>Value</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>------------------------</td>
<td>------------------------</td>
<td>----------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>kick_order</td>
<td>Optional</td>
<td>Increment used when adapting the basis order in function train methods.</td>
<td></td>
</tr>
<tr>
<td>max_order</td>
<td>Optional</td>
<td>Maximum polynomial order of each univariate function within the functional tensor train.</td>
<td></td>
</tr>
<tr>
<td>max_cv_order_candidates</td>
<td>Optional</td>
<td>Limit the number of cross-validation candidates for basis order.</td>
<td></td>
</tr>
<tr>
<td>start_rank</td>
<td>Optional</td>
<td>The initial rank used for the starting point during a rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>adapt_rank</td>
<td>Optional</td>
<td>Activate adaptive procedure for determining best rank representation.</td>
<td></td>
</tr>
<tr>
<td>kick_rank</td>
<td>Optional</td>
<td>The increment in rank employed during each iteration of the rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>max_rank</td>
<td>Optional</td>
<td>Limits the maximum rank that is explored during a rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>max_cv_rank_candidates</td>
<td>Optional</td>
<td>Limit the number of cross-validation candidates for rank.</td>
<td></td>
</tr>
<tr>
<td>samples_on_emulator</td>
<td>Optional</td>
<td>Number of samples at which to evaluate an emulator (surrogate).</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-------------</td>
<td>---------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
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</table>
### 7.2 method

<table>
<thead>
<tr>
<th>Optional (Choose One)</th>
<th>Covariance Type (Group 2)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate model values</td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

**Description**

Tensor train decompositions are approximations that exploit low rank structure in an input-output mapping. Refer to the function_train surrogate model for additional details.

**Usage Tips**

This method is a self-contained method alternative to the function_train surrogate model specification, similar to current method specifications for polynomial chaos and stochastic collocation. In particular, this function_train method specification directly couples with a simulation model (optionally identified with a model_pointer) and an additional function_train surrogate model specification is not required as these options have been embedded within the method specification.

**Examples**

```dakota
method, function_train
  start_order = 2
  start_rank = 2  kick_rank = 2  max_rank = 10
  adapt_rank
  solver_tolerance = 1e-12
  rounding_tolerance = 1e-12
  convergence_tol = 1e-6
  collocation_points = 100
  samples_on_emulator = 100000
  seed = 531
  response_levels = .1 1. 50. 100. 500. 1000.
  variance_based_decomp
```

**See Also**

These keywords may also be of interest:
Automatic polynomial order refinement

Specification

Alias: none

Argument(s): none

Default: no refinement

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
</tbody>
</table>

Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the `generalized` `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which
candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.48.1.1 uniform

- Keywords Area
- method
- function_train
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

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<th>Description of Group</th>
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<tr>
<td>Required (Choose One)</td>
<td>Uniform Refinement Approach (Group 1)</td>
<td>increment_start_rank</td>
<td>candidate generation by advancement of starting rank</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_start_order</td>
<td>candidate generation by advancement of starting basis order</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_max_rank</td>
<td>candidate generation by advancement of maximum rank</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_max_order</td>
<td>candidate generation by advancement of maximum basis order</td>
</tr>
</tbody>
</table>
Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.48.1.1.1 increment_start_rank

- Keywords Area
- method
- function_train
- p_refinement
- uniform
- increment_start_rank

candidate generation by advancement of starting rank

Specification

Alias: none

Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of start_rank to create new refinement candidates. It is typically used in the absence of adapt_rank to more rigidly define a refinement sequence.

7.2.48.1.1.2 increment_start_order

- Keywords Area
- method
- function_train
- p_refinement
- uniform
- increment_start_order

candidate generation by advancement of starting basis order
7.2 method

Specification

Alias: none
Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of \texttt{start\_order} to create new refinement candidates. It is typically used in the absence of \texttt{adapt\_order} to more rigidly define a refinement sequence.

7.2.48.1.1.3 \texttt{increment\_max\_rank}

- Keywords Area
- method
- function\_train
- \texttt{p\_refinement}
- uniform
- \texttt{increment\_max\_rank}

candidate generation by advancement of maximum rank

Specification

Alias: none
Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of \texttt{max\_rank} to create new refinement candidates. This option requires the use of \texttt{adapt\_rank}, for which this advancement creates an expanded range for the cross validation (CV) process. Once the CV process saturates (the best rank is less than the max), then no further advancement can occur and the refinement process (for a particular model level/fidelity) halts.

7.2.48.1.1.4 \texttt{increment\_max\_order}

- Keywords Area
- method
- function\_train
- \texttt{p\_refinement}
- uniform
- \texttt{increment\_max\_order}

candidate generation by advancement of maximum basis order
Specification

**Alias:** none  
**Argument(s):** none

**Description**

Within adaptive refinement processes, this specification selects the advancement of `max_order` to create new refinement candidates. This option requires the use of `adapt_order`, for which this advancement creates an expanded range for the cross validation (CV) process. Once the CV process saturates (the best order is less than the max), then no further advancement can occur and the refinement process (for a particular model level/fidelity) halts.

### 7.2.48.1.5 `increment_max_rank_order`

- **Keywords Area**
- **method**
- **function_train**
- **p_refinement**
- **uniform**
- **increment_max_rank_order**

Candidate generation by advancement of maximum rank and maximum basis order

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Within adaptive refinement processes, this specification selects the advancement of both `max_rank` and `max_order` to create new refinement candidates. This option requires the use of `adapt_rank` and `adapt_order`, for which this advancement creates expanded ranges for the cross validation (CV) process. Once the CV process saturates for one or more of these two dimensions (the best rank is less than the max rank and/or the best order is less than the max order), then no further advancement can occur in that dimension for a particular model level/fidelity.

### 7.2.48.2 `max_refinement_iterations`

- **Keywords Area**
- **method**
- **function_train**
- **max_refinement_iterations**

Maximum number of expansion refinement iterations
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

method, polynomial_chaos
dimension_adaptive generalized
  p_refinement
    max_refinement_iterations = 20
    convergence_tol = 1.e-4
  sparse_grid_level = 1

7.2.48.3 convergence_tolerance

• Keywords Area
  • method
  • function_train
  • convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library
This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

• DOT: relative tolerance that must be satisfied for two consecutive iterations

• NL2SOL: See nl2sol

• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance

• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.48.4 metric_scale

• Keywords Area

• method

• function_train

• metric_scale

define scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none

Argument(s): none

Default: relative

Child Keywords:

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<th>Description of Group scale of refinement metric (Group 1)</th>
<th>Dakota Keyword Description</th>
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<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
<td>Dakota Keyword</td>
</tr>
<tr>
<td></td>
<td>absolute</td>
<td>use of absolute statistical metrics when adapting UQ surrogates</td>
</tr>
</tbody>
</table>

Description

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.
7.2 method

Default Behavior
The default is relative.

Expected Output
Usage Tips
Additional Discussion

7.2.48.4.1 relative

• Keywords Area
• method
• function_train
• metric_scale
• relative

relative scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior
Expected Output
Usage Tips
Additional Discussion

7.2.48.4.2 absolute

• Keywords Area
• method
• function_train
• metric_scale
• absolute

use of absolute statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none
When adapting a UQ surrogate model, the absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.48.5 regression_type

- Keywords Area
- method
- function_train
- regression_type

Type of solver for forming function train approximations by regression

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td></td>
<td>ls</td>
<td>Use least squares solver for forming function train approximations by regression</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rls2</td>
<td>Use regularized regression solver for forming function train approximations</td>
</tr>
</tbody>
</table>

Description

Function train approximations are formed based on regression for the set of coefficients described at function_train. Solver options include least squares and regularized regression, where the latter penalizes high-order terms to mitigate overfitting.

Default Behavior

The default regression solver is least squares.

7.2.48.5.1 ls

- Keywords Area
Use least squares solver for forming function train approximations by regression

Specification

Alias: none
Argument(s): none

Description

Least squares is the standard solver for forming function train approximations based on regression. This option does not include regularization.

7.2.48.5.2 rls2

Use regularized regression solver for forming function train approximations

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>l2_penalty</td>
<td>Penalty value applied in regularized regression solver for function train approximations</td>
</tr>
</tbody>
</table>

Description

RLS2 is a regularized least squares solver that applies an $L2$ penalty to mitigate overfitting. It includes the option to specify a penalty parameter.
Penalty value applied in regularized regression solver for function train approximations

Specification

Alias: none

Argument(s): REAL

Description

Regularized regression applies a penalty to aspects of the recovered solution (e.g., cardinality, higher-order oscillatory terms, et al.) to mitigate overfitting. This specification provides the scalar penalty parameter (its value is not currently adapted or cross-validated).

Default Behavior

If this penalty parameter is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- rls2
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.48.7 max_cross_iterations

• Keywords Area
• method
• function_train
• max_cross_iterations

Maximum number of iterations for cross-approximation during a rank adaptation.

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The number of iterations for the cross-approximation algorithm is limited within the adapt_rank procedure.

See Also

These keywords may also be of interest:

• adapt_rank

7.2.48.8 solver_tolerance

• Keywords Area
• method
• function_train
• solver_tolerance

Convergence tolerance for the optimizer used during the regression solve.
Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0e-10

Description

Defines the function value and function gradient tolerance for optimizer used solving the regression problem for the functional tensor train coefficients.

7.2.48.9 response_scaling

- **Keywords Area**
- **method**
- **function_train**
- **response_scaling**

Perform bounds-scaling on response values prior to surrogate emulation

Specification

**Alias:** none

**Argument(s):** none

**Default:** use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.48.10 tensor_grid

- **Keywords Area**
- **method**
7.2 method

• function_train
• tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none
Default: regression with random sample set

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.48.11 collocation_points

• Keywords Area
• method
• function_train
• collocation_points

Number of collocation points used to estimate expansion coefficients

Specification

Alias: none
Argument(s): INTEGER

Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.
A corresponding sequence specification is documented at, e.g., collocation_points_sequence and collocation_points_sequence

See Also

These keywords may also be of interest:

• collocation_points_sequence method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence
7.2.48.12 collocation_ratio

- Keywords Area
- method
- function_train
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Description**

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2, produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^\alpha \) where \( c \) is the collocation_ratio and \( \alpha \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^\alpha n + 1 \)).

7.2.48.13 rounding_tolerance

- Keywords Area
- method
- function_train
- rounding_tolerance

An accuracy tolerance that is used to guide rounding during rank adaptation.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-10

**Description**

The rounding tolerance is used to measure how closely one can approximate a functional tensor train representation at a step of the optimization procedure using one of lower ranks. If this tolerance is small, then less compression is achieved but greater accuracy is obtained. If this tolerance is big, greater compression is achieved at the expense of accuracy.

**Default Behavior**

If this tolerance is not specified, then we adopt the internal C3 default.
See Also

These keywords may also be of interest:

- adapt_rank
- arithmetic_tolerance

7.2.48.14 arithmetic_tolerance

- Keywords Area
- method
- function_train
- arithmetic_tolerance

A secondary rounding tolerance used for post-processing

Specification

Alias: none

Argument(s): REAL

Default: 1.e-10

Description

After forming one or more function train approximations, this tolerance is used for rolling up final results, in particular, combining multilevel/multifidelity expansions and forming products for higher-order moment estimation. It is generally intended as a looser tolerance than rounding_tolerance.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- rounding_tolerance

7.2.48.15 start_order

- Keywords Area
- method
- function_train
- start_order

(Initial) polynomial order of each univariate function within the functional tensor train.
Specification

**Alias:** order

**Argument(s):** INTEGER

**Default:** 2

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
</tbody>
</table>

**Description**

This specifies the initial polynomial order of each univariate function that makes up the functional tensor train decomposition. In the case of `adapt_order`, polynomial order starts from this value and is then advanced; its maximum value is limited by `max_order` and increments are controlled by `kick_order`.

**Default Behavior**

Default is 2.

**See Also**

These keywords may also be of interest:

- `adapt_order`
- `kick_order`
- `max_order`

7.2.48.15.1 dimension_preference

- Keywords Area
- method
- function_train
- start_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids
Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.48.16 adapt_order

Activate adaptive procedure for determining the best basis order

Specification

Alias: none

Argument(s): none

Default: false

Description

This option activates a cross validation procedure to select the best basis order for representing the current set of data. It searches from start_order to max_order in increments of kick_order.

Default Behavior

No cross validation for basis order.

Examples

This example shows specification of an order adaptation starting at order 2, incrementing by 2, and limited at order 6.

model,
    id_model = 'FT'
    surrogate global function_train
    adapt_order start_order = 2 kick_order = 2 max_order = 6
    solver_tolerance = 1e-12
    rounding_tolerance = 1e-12
    dace_method_pointer = 'SAMPLING'

Note that adapt_rank and adapt_order can either be combined or used separately.
See Also

These keywords may also be of interest:

- start_order
- kick_order
- max_order

7.2.48.17 kick_order

- Keywords Area
- method
- function_train
- kick_order

increment used when adapting the basis order in function train methods

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

Used in combination with adapt_order, kick_order defines the size of the increment that is used during the cross validation process, within the range of start_order and max_order.

Default Behavior
Default is 1.

See Also

These keywords may also be of interest:

- adapt_order

7.2.48.18 max_order

- Keywords Area
- method
- function_train
- max_order

Maximum polynomial order of each univariate function within the functional tensor train.
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: unsigned short max

Description

This specification is inactive for current adaptation options.
In future adaptation options (i.e., cross-approximation adaptation), this will specify the maximum order that can be obtained during adaptation, where polynomial order will start from start_order and be limited by max_order.

Default Behavior
Max order is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- start_order

7.2.48.19 max_cv_order_candidates

- Keywords Area
- method
- function_train
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
The initial rank used for the starting point during a rank adaptation.

**Specification**

**Alias:** rank

**Argument(s):** INTEGER

**Default:** 2

**Description**

The _adapt_rank_ procedure increments the rank on each iteration, starting from the rank defined by `start_rank`

**Default Behavior**

Default is 2.

**See Also**

These keywords may also be of interest:

- _adapt_rank_
Specification

Alias: none
Argument(s): none
Default: false

Description

The adaptive algorithm proceeds as follows:

1. Start from rank \texttt{start\_rank} and form an approximation
2. Adapt the current approximation by searching for a solution with lower rank that achieves \texttt{L2} accuracy within epsilon tolerance of the reference.
3. If a lower rank solution is found with comparable accuracy, then stop. If not, increase the rank by an amount specified by \texttt{kick\_rank}.
4. Return to step 2 and continue until either \texttt{max\_rank} is reached or a converged rank (rank less than current reference with comparable accuracy) is found.

Default Behavior

No cross validation for rank.

Examples

This example shows specification of a rank adaptation starting at rank 2, incrementing by 2, and limited at rank 10.

```python
model,
    id_model = 'FT'
surrogate global function_train
    start_order = 5
    adapt_rank start_rank = 2 kick_rank = 2 max_rank = 10
    solver_tolerance = 1e-12
    rounding_tolerance = 1e-12
dace_method_pointer = 'SAMPLING'
```

Note that \texttt{adapt\_rank} and \texttt{adapt\_order} can either be combined or used separately.

See Also

These keywords may also be of interest:

- \texttt{start\_rank}
- \texttt{kick\_rank}
- \texttt{max\_rank}

The increment in rank employed during each iteration of the rank adaptation.
Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The `adapt_rank` procedure increments the rank on each iteration, using an increment defined by `kick_rank`

Default Behavior

Default is 1.

See Also

These keywords may also be of interest:

- `adapt_rank`

7.2.48.23 `max_rank`

- `Keywords Area`
- `method`
- `function_train`
- `max_rank`

Limits the maximum rank that is explored during a rank adaptation.

Specification

Alias: none
Argument(s): INTEGER
Default: `size_t max`

Description

The `adapt_rank` procedure increments the rank on each iteration, and the maximum value is limited by `max_rank`

Default Behavior

Max rank is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- `adapt_rank`
7.2.48.24  max_cv_rank_candidates

- Keywords Area
- method
- function_train
- max_cv_rank_candidates

Limit the number of cross-validation candidates for rank

Specification

Alias: none
Argument(s): INTEGER
Default: size_t max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_rank
- increment_max_rank_order
- function_train
- multifidelity_function_train

7.2.48.25  samples_on_emulator

- Keywords Area
- method
- function_train
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples
Argument(s): INTEGER
Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

**Default Behavior**

The default number of samples is method-dependent.

**Examples**

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method
  polynomial_chaos
    quadrature_order  = 2
    samples_on_emulator = 10000

7.2.48.26 sample_type
  • Keywords Area
  • method
  • function_train
  • sample_type

Selection of sampling strategy

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** random

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Type</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Sample Type (Group 1)</td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**
If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

Examples

```python
method
    sampling
        sample_type lhs
        samples = 20
        seed = 83921
```

7.2.48.26.1  lhs

- Keywords Area
- method
- function_train
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

**Alias:** none  
**Argument(s):** none

Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

```python
method
    sampling
        sample_type lhs
        samples = 20
```

7.2.48.26.2  random

- Keywords Area
- method
Uses purely random Monte Carlo sampling to sample variables

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The *random* keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the *random* keyword must be specified in conjunction with the *sample_type* keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```plaintext
method sampling
    sample_type random
    samples = 200

7.2.48.27 rng
```

**Selection of a random number generator**

**Specification**

**Alias:** none  
**Argument(s):** none

**Default:** Mersenne twister (mt19937)  
**Child Keywords:** none
Description

The `rng` keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

```
method
    sampling
        sample_type lhs
        samples = 10
        seed = 98765
        rng rnum2
```

7.2.48.27.1 mt19937

- **Keywords Area**
- **method**
- **function_train**
- **rng**
- **mt19937**

Generates random numbers using the Mersenne twister

Specification

**Alias:** none

**Argument(s):** none

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).
**Default Behavior**

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (mt19937) is recommended.

**Examples**

```dakota
method
   sampling
   sample_type lhs
   samples = 10
   seed = 98765
   rng mt19937
```

**rnum2**

Generates pseudo-random numbers using the Pecos package

**Specification**

*Alias:* none

*Argument(s):* none

**Description**

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (mt19937) is recommended over `rnum2`.

**Examples**

```dakota
method
   sampling
   sample_type lhs
   samples = 10
   seed = 98765
   rng rnum2
```
7.2.48.28 probability_refinement

- Keywords Area
- method
- function_train
- probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics
This keyword is related to the topics:
- reliability_methods

Specification

Alias: sample_refinement
Argument(s): none
Default: no refinement
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Importance Sampling Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design.</td>
<td></td>
</tr>
</tbody>
</table>

Description

The probability_refinement allows refinement of probability and generalized reliability results using importance sampling. If one specifies probability_refinement, there are some additional options. One can specify which type of importance sampling to use (import, adapt_import, or mm_adapt_import). Additionally, one can specify the number of refinement samples to use with refinement_samples and the seed to use with seed.

The probability_refinement density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.
7.2.48.28.1 import

- Keywords Area
- method
- function_train
- probability_refinement
- import

Importance sampling option for probability refinement

Specification

**Alias:** none

**Argument(s):** none

Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

7.2.48.28.2 adapt_import

- Keywords Area
- method
- function_train
- probability_refinement
- adapt_import

Importance sampling option for probability refinement

Specification

**Alias:** none

**Argument(s):** none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.
7.2.48.28.3  mm_adapt_import

- Keywords Area
- method
- function_train
- probability_refinement
- mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

`mm_adapt_import` starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then `mm_adapt_import` proceeds similarly to `adapt_import` (sample until convergence).

7.2.48.28.4  refinement_samples

- Keywords Area
- method
- function_train
- probability_refinement
- refinement_samples

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.
7.2.48.29 final_moments

- Keywords Area
- method
- function_train
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>none</td>
<td>final_moments</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

method, sampling
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments central

Overriding the default to none as follows:

method,
sampling
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments none

allows associated nested model mappings to be simplified from:

model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
        secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 1. 1. 0.
        secondary_response_mapping = 0. 0. 1.

7.2.48.29.1 none

• Keywords Area
• method
• function_train
• final_moments
• none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
sampling
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments none
7.2.48.29.2 standard

- Keywords Area
- method
- function_train
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```dakota
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments standard
```

7.2.48.29.3 central

- Keywords Area
- method
- function_train
- final_moments
- central

Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none
Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

7.2.48.30 response_levels

- Keywords Area
- method
- function_train
- response_levels

Values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakoata Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td>compute</td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior
If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_levels` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method, sampling, samples = 100 seed = 1
complementary_distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n', 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.

descriptors = 'TF1u', 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w', 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0

descriptors = 'TF1h', 'TF2h'
histogram_point_uncertain = 1
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
```
7.2 method

interface,
    system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses,
    response_functions = 3
    no_gradients
    no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td>5.0000000000e+01</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td>7.4000000000e+01</td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td>2.7000000000e+01</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td>6.1000000000e+01</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td>2.9000000000e+01</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td>9.0000000000e+01</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td>5.2000000000e+01</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td>9.0000000000e+01</td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.48.30.1 num_response_levels

- Keywords Area
- method
- function_train
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the `response_levels` will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```plaintext
method
sampling
  samples = 100
  seed = 34785
  num_response_levels = 1 1 1
  response_levels = 0.5 0.5 0.5
```

7.2.48.30.2 compute

- Keywords Area
- method
- function_train
- response_levels
- compute

Selection of statistics to compute at each response level
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td>system</td>
<td>system</td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

**Description**

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```
Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The **probabilities** keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If **response_levels** is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the **probabilities** keyword should be specified in conjunction with the **compute** keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```dakota
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```
Computes reliabilities associated with response levels

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `reliabilities` keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the reliabilities are not computed by default. To change this behavior, the `reliabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-reliability pairs according to the distribution defined.

**Examples**

```
method sampling
  sample_type = random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
  6.0e+04 6.5e+04 7.0e+04
  3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

Computes generalized reliabilities associated with response levels

**Specification**

**Alias:** none  
**Argument(s):** none
The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

**Expected Outputs**

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

**Examples**

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

7.2.48.30.2.4 system

- Keywords Area
- method
- function_train
- response_levels
- compute
- system

Compute system reliability (series or parallel)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>System Reliability Mode (Group 1)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.48.30.2.5 series

- Keywords Area
- method
- function_train
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.48.30.2.6 parallel

- Keywords Area
- method
- function_train
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system
Specification

**Alias:** none

**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.48.31 probability_levels

- Keywords Area
- method
- function_train
- probability_levels

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Examples**

The Dakota input file below specifies a sampling method with probability levels of interest.

Dakota Version 6.15 Reference Manual generated on November 5, 2021
method,
sampling,
samples = 100 seed = 1
complementary distribution
probability_levels = 1. .66 .33 0.
1. .8 .5 0.
1. .3 .2 0.

variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = ‘TF1n’ ‘TF2n’
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = ‘TF1u’ ‘TF2u’
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = ‘TF1w’ ‘TF2w’
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = ‘TF1h’ ‘TF2h’
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = ‘TF3h’

interface,
system async evaluation_concurrency = 5
analysis_driver = ‘text_book’

responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
Bin Lower Bin Upper Density Value
--------- --------- ------------
2.7604749078e+11 3.4221494996e+11 5.1384774972e-12
3.4221494996e+11 4.0634975300e+11 5.1454122311e-12
4.0634975300e+11 5.4196114379e+11 2.4334239039e-12

PDF for response_fn_2:
Bin Lower Bin Upper Density Value
--------- --------- ------------
4.6431154744e+04 5.651827775e+04 1.9839945149e-05
5.651827775e+04 6.1603813790e+04 5.8916108390e-05
6.1603813790e+04 7.8702465755e+04 2.9242071306e-05

PDF for response_fn_3:
Bin Lower Bin Upper Density Value
--------- --------- ------------
2.379673790e+05 3.6997214153e+05 5.3028386523e-06
3.6997214153e+05 3.8100966235e+05 9.0600055634e-06
3.8100966235e+05 4.4111498127e+05 3.3274925348e-06

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
Response Level Probability Level Reliability Index General Rel Index
-------------- ----------------- ----------------- ----------------- ------------------------------------
2.7604749078e+11 1.0000000000e+00
3.4221494996e+11 6.0000000000e-01

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.48.31.1 num_probability_levels

- Keywords Area
- method
- function_train
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.48.32 reliability_levels

- Keywords Area
- method
Specify reliability levels at which the response values will be estimated

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_reliability_levels</td>
<td>Specify which reliability_levels correspond to which response</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.48.32.1 num_reliability_levels

- **Keywords Area**
  - method
  - function_train
  - reliability_levels
  - num_reliability_levels

Specify which reliability_levels correspond to which response

**Specification**

**Alias:** none
**Argument(s):** INTEGERLIST  
**Default:** reliability_levels evenly distributed among response functions

**Description**
See parent page

7.2.48.33 gen_reliability_levels

- **Keywords Area**
- method
- function_train
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST  
**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_gen_reliability_levels</td>
<td></td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**
Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**
Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
7.2.48.33.1 num_gen_reliability_levels

- Keywords Area
- method
- function_train
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.48.34 distribution

- Keywords Area
- method
- function_train
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
</tbody>
</table>
The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**
If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**
Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```plaintext
distribution cumulative
```

Computes statistics according to cumulative functions
Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

method
    sampling
        sample_type lhs
        samples = 10
        distribution cumulative

7.2.48.34.2 complementary

- Keywords Area
- method
- function_train
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

method
    sampling
        sample_type lhs
        samples = 10
        distribution complementary

7.2.48.35 variance_based_decomp

- Keywords Area
• method
• function_train
• variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

Alias: none
Argument(s): none
Default: no variance-based decomposition

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>drop_tolerance</td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword variance_based_decomp. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

Default Behavior

Because of processing overhead and output volume, variance_based_decomp is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

Expected Outputs

When variance_based_decomp is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the interaction_order and drop_tolerance controls can further limit the computational and output requirements.
Examples

method, polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].

7.2.48.35.1 interaction_order

- Keywords Area
- method
- function_train
- variance_based_decomp
- interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

Specification

Alias: none

Argument(s): INTEGER

Default: Unrestricted (VBD includes all interaction orders present in the expansion)

Description

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume)

7.2.48.35.2 drop_tolerance

- Keywords Area
- method
- function_train
- variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance
Specification

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed

Description

The *drop_tolerance* keyword allows the user to specify a value below which sensitivity indices generated by *variance_based_decomp* are not displayed.

**Default Behavior**

By default, all sensitivity indices generated by *variance_based_decomp* are displayed.

**Usage Tips**

For *polynomial_chaos*, which outputs main, interaction, and total effects by default, the *univariate_effects* may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be *diagonal_covariance* or *full_covariance*, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

**Examples**

```yaml
method,
  sampling
    sample_type lhs
      samples = 100
    variance_based_decomp
      drop_tolerance = 0.001
```

7.2.48.36  *diagonal_covariance*

- Keywords Area
- method
- function_train
- *diagonal_covariance*

Display only the diagonal terms of the covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** *diagonal_covariance* for response vector > 10; else *full_covariance*

Description

With a large number of responses, the covariance matrix can be very large. *diagonal_covariance* is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
7.2.48.37  full covariance

- Keywords Area
- method
- function_train
- full covariance

Display the full covariance matrix

Specification

**Alias**: none

**Argument(s)**: none

Description

With a large number of responses, the covariance matrix can be very large. *full_covariance* is used to force Dakota to output the full covariance matrix.

7.2.48.38  import_approx_points_file

- Keywords Area
- method
- function_train
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

Specification

**Alias**: none

**Argument(s)**: **STRING**

**Default**: no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
**Description**

**Default Behavior**  No import of points at which to evaluate the surrogate.

**Expected Output**  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```plaintext
method
   polynomial_chaos
      expansion_order = 4
      import_approx_points_file = 'import.mcmc_annot.dat'
      annotated
```

7.2.48.38.1  custom_annotated

- Keywords Area
- method
- function_train
- import_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or
variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-
annotated allows user options for whether header row, eval_id column, and interface_id column appear
in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in
the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra
is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface-
_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.48.38.1.1 header

• Keywords Area

• method
• function_train
• import_approx_points_file
• custom_annotated
• header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.48.38.1.2 eval_id

• Keywords Area
• method
• function_train
• import_approx_points_file
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.48.38.1.3 interface_id

• Keywords Area
• method
• function_train
• import_approx_points_file
• custom_annotated
7.2 method

- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

**Description**

See description of parent custom_annotated

7.2.48.38.2 annotated

- Keywords Area
- method
- function_train
- import_approx_points_file
- annotated

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

**Usage Tips**

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.48.38.3 freeform

• Keywords Area

• method

• function_train

• import_approx_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format
7.2 method

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```dakota
environment
tabular_data
  tabular_data_file = ‘dakota_summary.dat’
  freeform

Resulting tabular file:

0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.48.38.4 active_only

• Keywords Area

• method

• function_train

• import_approx_points_file

• active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats
Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.48.39  export_approx_points_file

- Keywords Area
- method
- function_train
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file
Argument(s): STRING
Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td></td>
<td></td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td>freeform</td>
<td></td>
<td></td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:
7.2 method

- annotated (default)
- custom.annotated
- freeform

7.2.48.39.1 custom.annotated

- Keywords Area
- method
- function_train
- export_approx_points_file
- custom.annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.
Default Behavior

The *annotated* format is the default for tabular export/import. To control which header row and columns are in the input/output, specify `custom_annotated`, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to *annotated* format, though *freeform* remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface-_id), and data for variables and responses. Input file fragment:

```plaintext
evironment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   0.9  1.1  0.0002  0.26  0.76
2   0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.48.39.1.1 header

- Keywords Area
- method
- function_train
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias**: none

**Argument(s)**: none

**Default**: no header

Description

See description of parent `custom_annotated`
7.2.48.39.1.2 eval_id

- Keywords Area
- method
- function_train
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.48.39.1.3 interface_id

- Keywords Area
- method
- function_train
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated
7.2.48.39.2  annotated

- Keywords Area
- method
- function_train
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.48.39.3  freeform

- Keywords Area
- method
- function_train
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
```

7.2.48.40 seed

Keywords Area

- method
- function_train
- seed

Seed of the random number generator

Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.
7.2.48.41 fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.48.42 model_pointer

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

  • block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
    sample_type lhs
distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
single
  interface_pointer = 'I1'
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system async evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.49 multifidelity_function_train

• Keywords Area
• method
• multifidelity_function_train

Multifidelity uncertainty quantification using function train expansions

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
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<td></td>
<td>Maximum number of expansion refinement iterations</td>
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<td>Optional</td>
<td>convergence_tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
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<tr>
<td>Optional</td>
<td>metric_scale</td>
<td></td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td>statistics_mode</td>
<td>type of statistical metric roll-up for multifidelity UQ methods</td>
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</tr>
<tr>
<td>---------------</td>
<td>----------------</td>
<td>---------------------------------------------------------------</td>
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</tr>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
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</tr>
<tr>
<td>Optional</td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rounding_tolerance</td>
<td>An accuracy tolerance that is used to guide rounding during rank adaptation.</td>
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<tr>
<td>Optional</td>
<td>arithmetic_tolerance</td>
<td>A secondary rounding tolerance used for post-processing</td>
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<tr>
<td>Optional</td>
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<td>Type of solver for forming function train approximations by regression</td>
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</tr>
<tr>
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<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
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<td>Optional</td>
<td>max_cross_iterations</td>
<td>Maximum number of iterations for cross-approximation during a rank adaptation.</td>
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<td>Convergence tolerance for the optimizer used during the regression solve.</td>
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<tr>
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<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation</td>
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<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
<td></td>
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<tr>
<td>Optional</td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion</td>
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<td>start_order_sequence</td>
<td>Sequence of start orders used in a multi-stage expansion</td>
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</tr>
<tr>
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<td>adapt_order</td>
<td>Activate adaptive procedure for determining the best basis order</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>kick_order</td>
<td>increment used when adapting the basis order in function train methods</td>
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<tr>
<td>Optional</td>
<td>max_order</td>
<td>Maximum polynomial order of each univariate function within the functional tensor train</td>
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<tr>
<td>Optional</td>
<td>max_cv_order_ - candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
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<tr>
<td>Optional</td>
<td>start_rank_ - sequence</td>
<td>Sequence of start ranks used in a multi-stage expansion</td>
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<tr>
<td>Optional</td>
<td>adapt_rank</td>
<td>Activate adaptive procedure for determining best rank representation</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>kick_rank</td>
<td>The increment in rank employed during each iteration of the rank adaptation.</td>
<td></td>
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<tr>
<td>Optional</td>
<td>max_rank</td>
<td>Limits the maximum rank that is explored during a rank adaptation.</td>
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</tr>
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<td>Optional</td>
<td>max_cv_rank_ - candidates</td>
<td>Limit the number of cross-validation candidates for rank</td>
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<td>samples_on_ - emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
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<td>sample_type</td>
<td>Selection of sampling strategy</td>
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<td>rng</td>
<td>Selection of a random number generator</td>
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<td>probability_ - refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
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<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
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<tr>
<td>-------------------</td>
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<td>------------------------------------------------------------------------------------------</td>
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<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
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<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
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<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
</tr>
<tr>
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<td>Covariance Type (Group 1)</td>
<td>diagonal_covariance Display only the diagonal terms of the covariance matrix</td>
<td></td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate</td>
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### Keywords Area

<table>
<thead>
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<th>Optional</th>
<th>export_approx_points_file</th>
<th>Output file for surrogate model value evaluations</th>
</tr>
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<td>Optional</td>
<td>seed_sequence</td>
<td>Sequence of seed values for multi-stage random sampling</td>
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<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
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<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
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</table>

### Description

As described in the `function_train` method and the `function_train` model, the function train (FT) approximation is a polynomial expansion that exploits low rank structure within the mapping from input random variables to output quantities of interest (QoI). For multilevel and multifidelity function train approximations, we decompose this expansion into several constituent expansions, one per model form or solution control level, where independent function train approximations are constructed for the low-fidelity/coarse resolution model and one or more levels of model discrepancy.

In a three-model case with low-fidelity (L), medium-fidelity (M), and high-fidelity (H) models and an additive discrepancy approach, we can denote this as:

\[
Q^H \approx \hat{Q}^L + \hat{\Delta}^{ML} + \hat{\Delta}^{HM}
\]

where \( \Delta^{ij} \) represents a discrepancy expansion computed from \( Q^i - Q^j \) and reduced rank representations of these discrepancies may be targeted (\( r_{HM} < r_{ML} < r_L \)).

In multifidelity approaches, sample allocation for the constituent expansions can be performed with either no, individual, or integrated adaptive refinement as described in `allocation_control`.

### Expected HDF5 Output

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments (expansion moments only)
- Probability Density
- Level Mappings

In addition, the execution group has the attribute `equiv_hf_evals`, which records the equivalent number of high-fidelity evaluations.

### See Also

These keywords may also be of interest:

- `function_train`
- `function_train`
7.2.49.1  p_refinement

- Keywords Area
  - method
  - multifidelity_function_train
  - p_refinement

Automatic polynomial order refinement

Specification

Alias: none
Argument(s): none
Default: no refinement
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
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<tr>
<td>Required</td>
<td></td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
</tbody>
</table>

Description

The p_refinement keyword specifies the usage of automated polynomial order refinement, which can be either uniform or dimension_adaptive.

The dimension_adaptive option is supported for the tensor-product quadrature and Smolyak sparse grid options and uniform is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the max_iterations and convergence_tolerance method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The dimension_adaptive case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the uniform refinement case, the quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
7.2.49.1.1 uniform

- Keywords Area
- method
- multifidelity_function_train
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

Specification

Alias: none
Argument(s): none

Child Keywords:

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<th>Dakota Keyword Description</th>
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<td>increment_start_-rank</td>
<td>candidate generation by advancement of starting rank</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_start_-order</td>
<td>candidate generation by advancement of starting basis order</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_max_-rank</td>
<td>candidate generation by advancement of maximum rank</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_max_-order</td>
<td>candidate generation by advancement of maximum basis order</td>
</tr>
<tr>
<td></td>
<td></td>
<td>increment_max_-rank_order</td>
<td>candidate generation by advancement of maximum rank and maximum basis order</td>
</tr>
</tbody>
</table>

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_-order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.
### 7.2 method

#### 7.2.49.1.1.1 increment_start_rank

- **Keywords Area**
  - method
  - multifidelity_function_train
  - p_refinement
  - uniform
  - increment_start_rank

Candidate generation by advancement of starting rank

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Within adaptive refinement processes, this specification selects the advancement of `start_rank` to create new refinement candidates. It is typically used in the absence of `adapt_rank` to more rigidly define a refinement sequence.

#### 7.2.49.1.1.2 increment_start_order

- **Keywords Area**
  - method
  - multifidelity_function_train
  - p_refinement
  - uniform
  - increment_start_order

Candidate generation by advancement of starting basis order

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Within adaptive refinement processes, this specification selects the advancement of `start_order` to create new refinement candidates. It is typically used in the absence of `adapt_order` to more rigidly define a refinement sequence.
7.2.49.1.1.3 increment_max_rank

- Keywords Area
- method
- multifidelity_function_train
- p_refinement
- uniform
- increment_max_rank

candidate generation by advancement of maximum rank

Specification

Alias: none
Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of max_rank to create new refinement candidates. This option requires the use of adapt_rank, for which this advancement creates an expanded range for the cross validation (CV) process. Once the CV process saturates (the best rank is less than the max), then no further advancement can occur and the refinement process (for a particular model level/fidelity) halts.

7.2.49.1.1.4 increment_max_order

- Keywords Area
- method
- multifidelity_function_train
- p_refinement
- uniform
- increment_max_order

candidate generation by advancement of maximum basis order

Specification

Alias: none
Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of max_order to create new refinement candidates. This option requires the use of adapt_order, for which this advancement creates an expanded range for the cross validation (CV) process. Once the CV process saturates (the best order is less than the max), then no further advancement can occur and the refinement process (for a particular model level/fidelity) halts.
7.2.49.1.5 increment_max_rank_order

• Keywords Area
• method
• multifidelity_function_train
• p_refinement
• uniform
• increment_max_rank_order

candidate generation by advancement of maximum rank and maximum basis order

Specification

Alias: none
Argument(s): none

Description

Within adaptive refinement processes, this specification selects the advancement of both max_rank and max_order to create new refinement candidates. This option requires the use of adapt_rank and adapt_order, for which this advancement creates expanded ranges for the cross validation (CV) process. Once the CV process saturates for one or more of these two dimensions (the best rank is less than the max rank and/or the best order is less than the max order), then no further advancement can occur in that dimension for a particular model level/fidelity.

7.2.49.2 max_refinement_iterations

• Keywords Area
• method
• multifidelity_function_train
• max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

```plaintext
method,  
  polynomial_chaos  
  dimension_adaptive generalized  
  p_refinement  
    max_refinement_iterations = 20  
    convergence_tol = 1.e-4  
  sparse_grid_level = 1
```

7.2.49.3 convergence_tolerance

- Keywords Area
- method
- multifidelity_function_train
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration.

For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.49.4 metric_scale

• Keywords Area
• method
• multifidelity_function_train
• metric_scale

define scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none
Default: relative
Child Keywords:

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<th>Description of Group scale of refinement metric (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>relative</td>
<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>absolute</td>
<td>use of absolute statistical metrics when adapting UQ surrogates</td>
</tr>
</tbody>
</table>

Description

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior

The default is relative.

Expected Output

Usage Tips

Additional Discussion

7.2.49.4.1 relative

• Keywords Area
• method
relative scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.49.4.2 absolute

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

Default Behavior

Expected Output

Usage Tips

Additional Discussion
7.2.49.5 statistics_mode

- Keywords Area
- method
- multifidelity_function_train
- statistics_mode

type of statistical metric roll-up for multifidelity UQ methods

Specification

**Alias:** none

**Argument(s):** none

**Default:** DEFAULT_EXPANSION_STATS

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>active</td>
<td>use statistics for the active expansion within multifidelity refinement</td>
<td>active</td>
<td></td>
</tr>
<tr>
<td>combined</td>
<td>use statistics for the combined expansion within multifidelity refinement</td>
<td>combined</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

When performing adaptive refinement in multifidelity UQ methods, one can adapt the expansion for a model level/fidelity based on its local statistics (active) or based on the statistics for the aggregate expansion (combined). The latter case may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

**Default Behavior**

Default for multifidelity methods is combined statistics.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.49.5.1 active

- Keywords Area
- method
- multifidelity_function_train
- statistics_mode
use statistics for the active expansion within multifidelity refinement

Specification

Alias: none
Argument(s): none

Description

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the active model level/fidelity. This approach avoids some computational overhead in some cases by avoiding expansion roll-up.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

use statistics for the combined expansion within multifidelity refinement

Specification

Alias: none
Argument(s): none

Description

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the aggregate multifidelity expansion. This approach may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

Default Behavior

Expected Output

Usage Tips

Additional Discussion
7.2.49.6 allocation_control

- Keywords Area
- method
- multifidelity_function_train
- allocation_control

Sample allocation approach for multifidelity expansions

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Multifidelity Sample Allocation Control (Group 1)</td>
<td>greedy</td>
<td>Sample allocation based on greedy refinement within multifidelity function train</td>
</tr>
</tbody>
</table>

**Description**

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

**Default Behavior**

The default, when allocation_control is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

7.2.49.6.1 greedy

- Keywords Area
- method
- multifidelity_function_train
- allocation_control
- greedy

Sample allocation based on greedy refinement within multifidelity function train

**Specification**

**Alias:** none  
**Argument(s):** none
Description

Multifidelity function train supports greedy refinement strategies based on regression approaches for computing expansion coefficients. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within a unified competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any $\beta^*/\beta/\beta$ level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.

Examples

The following example of greedy multifidelity function train starts from a rank-two order-two reference expansion for each level, with twice as many samples as regression coefficients, and generates candidate refinements for each level that are competed in an integrated greedy competition. The number of new samples for the incremented candidate is determined from the collocation ratio times the regression size (which may either be fixed or adapted in the case of adapt_rank). In this example, the number of candidates for each level is limited to one uniform refinement of the current expansion, and uniform refinement currently involves an advancement in the basis order for all approximation cores in combination with a rank adaptation between two and ten, incrementing in steps of two.

```plaintext
method,
    model_pointer = 'HIERARCH'
    multifidelity_function_train
    allocation_control greedy
    p_refinement uniform
    start_rank_sequence = 2 2 2 2
    adapt_rank kick_rank = 2 max_rank = 10
    start_order_sequence = 2 2 2 2 max_order = 10
    collocation_ratio = 2. seed = 160415
    convergence_tolerance = 1.e-2
    max_refinement_iterations = 5
```

7.2.49.7 discrepancy_emulation

- Keywords Area
- method
- multifidelity_function_train
- discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none
Default: distinct
Child Keywords:
In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.

### Examples

```plaintext
method, multilevel_polynomial_chaos
  expansion_order_sequence = 2
  collocation_ratio = .9
  orthogonal_matching_pursuit
  discrepancy_emulation recursive
```

**7.2.49.7.1 distinct**

- **Keywords Area**
- **method**
  - **multifidelity_function_train**
  - **discrepancy_emulation**
  - **distinct**

Distinct formulation for emulation of model discrepancies.

### Specification

**Alias:** paired

**Argument(s):** none
Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.49.7.2 recursive

- Keywords Area
- method
- multifidelity_function_train
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.49.8 rounding_tolerance

- Keywords Area
- method
- multifidelity_function_train
- rounding_tolerance

An accuracy tolerance that is used to guide rounding during rank adaptation.

Specification

Alias: none
Argument(s): REAL
Default: 1.e-10
Description

The rounding tolerance is used to measure how closely one can approximate a functional tensor train representation at a step of the optimization procedure using one of lower ranks. If this tolerance is small, then less compression is achieved but greater accuracy is obtained. If this tolerance is big, greater compression is achieved at the expense of accuracy.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- adapt_rank
- arithmetic_tolerance

7.2.49.9 arithmetic_tolerance

A secondary rounding tolerance used for post-processing

Specification

Alias: none
Argument(s): REAL
Default: 1.e-10

Description

After forming one or more function train approximations, this tolerance is used for rolling up final results, in particular, combining multilevel/multifidelity expansions and forming products for higher-order moment estimation. It is generally intended as a looser tolerance than rounding_tolerance.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- rounding_tolerance
7.2.49.10 regression_type

- Keywords Area
- method
- multifidelity_function_train
- regression_type

Type of solver for forming function train approximations by regression

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<td></td>
<td></td>
<td>ls</td>
<td>Use least squares solver for forming function train approximations by regression</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rls2</td>
<td>Use regularized regression solver for forming function train approximations</td>
</tr>
</tbody>
</table>

**Description**

Function train approximations are formed based on regression for the set of coefficients described at function_train. Solver options include least squares and regularized regression, where the latter penalizes high-order terms to mitigate overfitting.

**Default Behavior**

The default regression solver is least squares.

7.2.49.10.1 ls

- Keywords Area
- method
- multifidelity_function_train
- regression_type
- ls

Use least squares solver for forming function train approximations by regression
7.2 method

Specification

**Alias:** none

**Argument(s):** none

Description

Least squares is the standard solver for forming function train approximations based on regression. This option does not include regularization.

7.2.49.10.2 rls2

- Keywords Area
- method
- multifidelity_function_train
- regression_type
- rls2

Use regularized regression solver for forming function train approximations

Specification

**Alias:** none

**Argument(s):** none

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>l2_penalty</td>
<td>Penalty value applied in regularized regression solver for function train approximations</td>
</tr>
</tbody>
</table>

Description

RLS2 is a regularized least squares solver that applies an $L2$ penalty to mitigate overfitting. It includes the option to specify a penalty parameter.

See Also

These keywords may also be of interest:

- l2_penalty

7.2.49.10.2.1 l2_penalty

- Keywords Area
- method
• multifidelity_function_train
• regression_type
• rls2
• l2_penalty

Penalty value applied in regularized regression solver for function train approximations

Specification

Alias: none
Argument(s): REAL

Description

Regularized regression applies a penalty to aspects of the recovered solution (e.g., cardinality, higher-order oscillatory terms, et al.) to mitigate overfitting. This specification provides the scalar penalty parameter (its value is not currently adapted or cross-validated).

Default Behavior

If this penalty parameter is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

• rls2

7.2.49.11 max_solver_iterations

• Keywords Area
• method
• multifidelity_function_train
• max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.
7.2.49.12 max_cross_iterations

- Keywords Area
- method
- multifidelity_function_train
- max_cross_iterations

Maximum number of iterations for cross-approximation during a rank adaptation.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1

**Description**

The number of iterations for the cross-approximation algorithm is limited within the `adapt_rank` procedure.

**See Also**

These keywords may also be of interest:

- `adapt_rank`

7.2.49.13 solver_tolerance

- Keywords Area
- method
- multifidelity_function_train
- solver_tolerance

Convergence tolerance for the optimizer used during the regression solve.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0e-10

**Description**

Defines the function value and function gradient tolerance for optimizer used solving the regression problem for the functional tensor train coefficients.
7.2.49.14 response_scaling

- Keywords Area
- method
- multifidelity_function_train
- response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.49.15 tensor_grid

- Keywords Area
- method
- multifidelity_function_train
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with random sample set
Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.49.16 collocation_points_sequence

- Keywords Area
- method
- multifidelity_function_train
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

Specification

Alias: pilot_samples

Argument(s): INTEGERLIST

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.49.17 collocation_ratio

- Keywords Area
- method
- multifidelity_function_train
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.
Specification

Alias: none
Argument(s): REAL
Default: 0.

Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2, produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o \)).

7.2.49.18 start_order_sequence

- Keywords Area
- method
- multifidelity_function_train
- start_order_sequence

Sequence of start orders used in a multi-stage expansion

Specification

Alias: order_sequence
Argument(s): INTEGERLIST
Default: 2

Child Keywords:

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<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
</tbody>
</table>

Description

Each level entry of the start_order_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support start order sequences include multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., start_order
See Also

These keywords may also be of interest:

- start_order

7.2.49.18.1 dimension_preference

- Keywords Area
- method
- multifidelity_function_trian
- start_order_sequence
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.49.19 adapt_order

- Keywords Area
- method
- multifidelity_function_train
- adapt_order

Activate adaptive procedure for determining the best basis order

Specification

**Alias:** none

**Argument(s):** none

**Default:** false
Description

This option activates a cross validation procedure to select the best basis order for representing the current set of data. It searches from \texttt{start\_order} to \texttt{max\_order} in increments of \texttt{kick\_order}.

Default Behavior

No cross validation for basis order.

Examples

This example shows specification of an order adaptation starting at order 2, incrementing by 2, and limited at order 6.

```plaintext
model, id_model = 'FT'
surrogate global function_train
  start\_rank = 4
  adapt\_order start\_order = 2 kick\_order = 2 max\_order = 6
  solver\_tolerance = 1e-12
  rounding\_tolerance = 1e-12
dace\_method\_pointer = 'SAMPLING'
```

Note that \texttt{adapt\_rank} and \texttt{adapt\_order} can either be combined or used separately.

See Also

These keywords may also be of interest:

- \texttt{start\_order}
- \texttt{kick\_order}
- \texttt{max\_order}

7.2.49.20 \texttt{kick\_order}

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{multifidelity\_function\_train}
- \texttt{kick\_order}

increment used when adapting the basis order in function train methods

Specification

\textbf{Alias:} none

\textbf{Argument(s):} INTEGER

\textbf{Default:} 1
Description

Used in combination with adapt_order, kick_order defines the size of the increment that is used during the cross validation process, within the range of start_order and max_order.

Default Behavior
Default is 1.

See Also

These keywords may also be of interest:

- adapt_order

7.2.49.21 max_order

- Keywords Area
- method
- multifidelity_function_train
- max_order

Maximum polynomial order of each univariate function within the functional tensor train.

Specification

Alias: none
Argument(s): INTEGER
Default: unsigned short max

Description

This specification is inactive for current adaptation options.
In future adaptation options (i.e., cross-approximation adaptation), this will specify the maximum order that can be obtained during adaptation, where polynomial order will start from start_order and be limited by max_order.

Default Behavior
Max order is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- start_order

7.2.49.22 max_cv_order_candidates

- Keywords Area
- method
Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** ushort max

**Description**

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

**See Also**

These keywords may also be of interest:

- `increment_max_order`
- `increment_max_rank_order`
- `polynomial_chaos`
- `multifidelity_polynomial_chaos`
- `function_train`
- `multifidelity_function_train`

**7.2.49.23 start_rank_sequence**

**Specification**

**Alias:** rank_sequence  
**Argument(s):** INTEGERLIST  
**Default:** 2

Sequence of start ranks used in a multi-stage expansion
Description

Each level entry of the `start_rank_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support start rank sequences include multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `start_rank`

See Also

These keywords may also be of interest:

- `start_rank`

7.2.49.24 adapt_rank

- Keywords Area
- method
- multifidelity_function_train
- adapt_rank

Activate adaptive procedure for determining best rank representation

Specification

**Alias:** none

**Argument(s):** none

**Default:** false

Description

The adaptive algorithm proceeds as follows:

1. Start from rank `start_rank` and form an approximation

2. Adapt the current approximation by searching for a solution with lower rank that achieves L2 accuracy within epsilon tolerance of the reference.

3. If a lower rank solution is found with comparable accuracy, then stop. If not, increase the rank by an amount specified by `kick_rank`.

4. Return to step 2 and continue until either `max_rank` is reached or a converged rank (rank less than current reference with comparable accuracy) is found.

Default Behavior

No cross validation for rank.
Examples

This example shows specification of a rank adaptation starting at rank 2, incrementing by 2, and limited at rank 10.

```dakota
model,
    id_model = 'FT'
    surrogate global function_train
    start_order = 5
    adapt_rank start_rank = 2 kick_rank = 2 max_rank = 10
    solver_tolerance = 1e-12
    rounding_tolerance = 1e-12
    dace_method_pointer = 'SAMPLING'
```

Note that adapt_rank and adapt_order can either be combined or used separately.

See Also

These keywords may also be of interest:

- start_rank
- kick_rank
- max_rank

7.2.49.25 kick_rank

- Keywords Area
- method
- multifidelity_function_train
- kick_rank

The increment in rank employed during each iteration of the rank adaptation.

Specification

Alias: none

Argument(s): INTEGER

Default: 1

Description

The adapt_rank procedure increments the rank on each iteration, using an increment defined by kick_rank.

Default Behavior

Default is 1.

See Also

These keywords may also be of interest:

- adapt_rank
7.2.49.26  max_rank

- Keywords Area
- method
- multifidelity_function_train
- max_rank

Limits the maximum rank that is explored during a rank adaptation.

Specification

Alias: none

Argument(s): INTEGER

Default: size_t max

Description

The adapt_rank procedure increments the rank on each iteration, and the maximum value is limited by max_rank

Default Behavior

Max rank is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- adapt_rank

7.2.49.27  max_cv_rank_candidates

- Keywords Area
- method
- multifidelity_function_train
- max_cv_rank_candidates

Limit the number of cross-validation candidates for rank

Specification

Alias: none

Argument(s): INTEGER

Default: size_t max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_rank
- increment_max_rank_order
- function_train
- multifidelity_function_train

7.2.49.28 samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

**Alias:** samples

**Argument(s):** INTEGER

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

**Default Behavior**

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
  polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```
The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921

7.2.49.29.1 (lhs)
```

• sample_type

• lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

Alias: none

Argument(s): none

Description

The lhs keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the lhs keyword must appear in conjunction with the sample_type keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

method
  sampling
    sample_type lhs
    samples = 20

7.2.49.29.2 random

  • Keywords Area

  • method

  • multifidelity_function_train

  • sample_type

  • random

Uses purely random Monte Carlo sampling to sample variables

Specification

Alias: none

Argument(s): none
The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior
In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

Usage Tips
Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```plaintext
method sampling
  sample_type random
  samples = 200
```

7.2.49.30 rng

- Keywords Area
- method
- multifidelity_function_train
- rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none
Default: Mersenne twister (mt19937)

Child Keywords:

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<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

Description
The rng keyword is used to indicate a choice of random number generator.

Default Behavior
If specified, the **rng** keyword must be accompanied by either **rnum2** (pseudo-random numbers) or **mt19937** (random numbers generated by the Mersenne twister). Otherwise, **mt19937**, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

```
7.2.49.30.1  mt19937
  • Keywords Area
  • method
  • multifidelity_function_train
  • rng
  • mt19937
```

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The **mt19937** keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

**Default Behavior**

**mt19937** is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the **rng** keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (**mt19937**) is recommended.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```
7.2.49.30.2  rnum2

- Keywords Area
- method
- multifidelity_function_train
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

**Alias:** none

**Argument(s):** none

Description

The \texttt{rnum2} keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

\texttt{rnum2} is not used by default. To change this behavior, it must be specified in conjunction with the \texttt{rng} keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended over \texttt{rnum2}.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.49.31  probability_refinement

- Keywords Area
- method
- multifidelity_function_train
- probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

- reliability_methods
### Specification

**Alias:** sample_refinement  
**Argument(s):** none  
**Default:** no refinement  

### Child Keywords:

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<th>Dakota Keyword</th>
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<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
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<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>

| | | Optional | Number of samples used to refine a probability estimate or sampling design. |

### Description

The `probability_refinement` allows refinement of probability and generalized reliability results using importance sampling. If one specifies `probability_refinement`, there are some additional options. One can specify which type of importance sampling to use (`import`, `adapt_import`, or `mm_adapt_import`). Additionally, one can specify the number of refinement samples to use with `refinement_samples` and the seed to use with `seed`.

The `probability_refinement` density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

### 7.2.49.31.1 import

- Keywords Area
- method
- multifidelity_function_train
- probability_refinement
- import

Importance sampling option for probability refinement

### Specification

**Alias:** none  
**Argument(s):** none
Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then gener-
ates the importance samples, weights them by their probability of occurrence given the original density, and calcu-
lates the required probability (CDF or CCDF level).

7.2.49.31.2  adapt_import

- Keywords Area
- method
- multifidelity_function_train
- probability_refinement
- adapt_import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It
then generates the importance samples, weights them by their probability of occurrence given the original density,
and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability
estimate converges.

7.2.49.31.3  mm_adapt_import

- Keywords Area
- method
- multifidelity_function_train
- probability_refinement
- mm_adapt_import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling
density. First, it uses a small number of samples around each of the initial samples in the failure region. Note
that these samples are allocated to the different points based on their relative probabilities of occurrence: more
probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then \texttt{mm\_adapt\_import} proceeds similarly to \texttt{adapt\_import} (sample until convergence).

7.2.49.31.4 refinement_samples

- Keywords Area
- method
- multifidelity\_function\_train
- probability\_refinement
- refinement\_samples

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using uni-modal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial\_samples in a sampling design.

7.2.49.32 final_moments

- Keywords Area
- method
- multifidelity\_function\_train
- final\_moments

Output moments of the specified type and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** standard

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
### Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

### Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
        secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 1. 1. 0.
        secondary_response_mapping = 0. 0. 1.
```
7.2.49.32.1 none

- Keywords Area
- method
- multifidelity_function_train
- final_moments
- none

Omit moments from the set of final statistics.

**Specification**

Alias: none

Argument(s): none

**Description**

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

**Examples**

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

7.2.49.32.2 standard

- Keywords Area
- method
- multifidelity_function_train
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

**Specification**

Alias: none

Argument(s): none

**Description**

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.
Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50  seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments standard
```

7.2.49.32.3  central

- Keywords Area
- method
- multifidelity_function_train
- final_moments
- central

Output central moments and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50  seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

7.2.49.33  response_levels

- Keywords Area
- method
- multifidelity_function_train
- response_levels

Values at which to estimate desired statistics for each response
Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

Examples

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num-
The Dakota input file below specifies a sampling method with response levels of interest.

method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05

variables,

normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = ’TF1n’ ’TF2n’

uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = ’TF1u’ ’TF2u’

weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = ’TF1w’ ’TF2w’

histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = ’TF1h’ ’TF2h’

histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = ’TF3h’

interface,
system asynch evaluation_concurrency = 5
analysis_driver = ’text_book’

responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643115474e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>1.0341896485e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
</tbody>
</table>
Level mappings for each response function:

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.49.33.1 num_response_levels

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.
Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.49.33.2 compute

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute

Selection of statistics to compute at each response level

Specification

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
</tbody>
</table>
The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

**7.2.49.33.2.1 probabilities**

- **Keywords Area**
  - `method`
  - `multifidelity_function_train`
  - `response_levels`
  - `compute`
  - `probabilities`

Computes probabilities associated with response levels

**Specification**

- **Alias**: none
- **Argument(s)**: none
Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

7.2.49.33.2.2  reliabilities

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute
- reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The `reliabilities` keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the reliabilities are not computed by default. To change this behavior, the `reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

7.2.49.33.2.3 gen_reliabilities

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```
7.2.49.33.2.4 system

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute
- system

Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td>parallel</td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.49.33.2.5 series

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Specification

**Alias:** none
**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.49.33.2.6 parallel

- Keywords Area
- method
- multifidelity_function_train
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

**Alias:** none
**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.49.34 probability_levels

- Keywords Area
- method
- multifidelity_function_train
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

**Alias:** none
**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**
### 7.2 Method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

#### Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

#### Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

#### Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```plaintext
method, 
  sampling,
  samples = 100 seed = 1 
  complementary distribution 
  probability_levels = 1. .66 .33 0. 
  1. .8 .5 0. 
  1. .3 .2 0. 
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond to the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.699721453e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.699721453e+05</td>
<td>3.810096235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.810096235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.699721453e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.810096235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.49.34.1 num_probability_levels

- Keywords Area
- method
• multifidelity_function_train
• probability_levels
• num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none
Argument(s): INTEGERLIST
Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.49.35 reliability_levels

• Keywords Area
• method
• multifidelity_function_train
• reliability_levels

Specify reliability levels at which the response values will be estimated

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td>none</td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.49.35.1 num_reliability_levels

• Keywords Area
• method
• multifidelity_function_train
• reliability_levels
• num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.49.36 gen_reliability_levels

• Keywords Area
• method
• multifidelity_function_train
• gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.49.36.1 num_gen_reliability_levels

- Keywords Area
- method
- multifidelity_function_train
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.49.37 distribution

- Keywords Area
- method
Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
  distribution cumulative
```

**7.2.49.37.1 cumulative**

- **Keywords Area**
- **method**
- **multifidelity_function_train**
7.2 method

- distribution
- cumulative

Computes statistics according to cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the **cumulative** keyword must appear in conjunction with the **distribution** keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```plaintext
method
    sampling
        sample_type lhs
        samples = 10
        distribution cumulative
```

7.2.49.37.2 complementary

- Keywords Area
- method
- multifidelity_function_train
- distribution
- complementary

Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```
method
    sampling
        sample_type lhs
        samples = 10
        distribution complementary
```

7.2.49.38 variance_based_decomp

- Keywords Area
- method
- multifidelity_function_train
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

Alias: none

Argument(s): none

Default: no variance-based decomposition

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics. Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>drop_tolerance</td>
<td></td>
</tr>
</tbody>
</table>
Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword \texttt{variance-based_decomp}. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

Default Behavior

Because of processing overhead and output volume, \texttt{variance-based_decomp} is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

Expected Outputs

When \texttt{variance-based_decomp} is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of \textit{univariate} terms for each individual variable, interaction effects represent the set of \textit{mixed} terms (the complement of the univariate set), and total effects represent the \textit{complete} set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the \texttt{interaction_order} and \texttt{drop_tolerance} controls can further limit the computational and output requirements.

Examples

\begin{verbatim}
  method,  
  polynomial_chaos # or stoch_collocation  
  sparse_grid_level = 3  
  variance_based_decomp interaction_order = 2
\end{verbatim}

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of \cite{76}: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in \cite{90}.

7.2.49.38.1 interaction_order

- Keywords Area
  - method
    - multifidelity_function_train
    - variance_based_decomp
    - interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

Description

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.49.38.2 drop_tolerance

- **Keywords Area**
- **method**
- **multifidelity_function_train**
- **variance_based_decomp**
- **drop_tolerance**

Suppresses output of sensitivity indices with values lower than this tolerance

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed

Description

The `drop_tolerance` keyword allows the user to specify a value below which sensitivity indices generated by `variance_based_decomp` are not displayed.

Default Behavior

By default, all sensitivity indices generated by `variance_based_decomp` are displayed.

Usage Tips

For `polynomial_chaos`, which outputs main, interaction, and total effects by default, the `univariate_effects` may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

method,
7.2 method

sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
  drop_tolerance = 0.001

7.2.49.39 diagonal_covariance

- Keywords Area
- method
- multifidelity_function_train
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none
Argument(s): none
Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.49.40 full_covariance

- Keywords Area
- method
- multifidelity_function_train
- full_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.
7.2.49.41 import_approx_points_file

- Keywords Area
- method
- multifidelity_function_train
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>annotated</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td></td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

**Default Behavior** No import of points at which to evaluate the surrogate.

**Expected Output** The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```dakota
method
  polynomial_chaos
    expansion_order = 4
    import_approx_points_file = 'import.mcmc_annot.dat'
    annotated
```

7.2.49.41.1 custom_annotated

- Keywords Area
- method
- multifidelity_function_train
• import_approx_points_file
  • custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:
  • file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom.annotated header eval_id
```

Resulting tabular file:

```
%eval_id   x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1   0.9   1.1   0.0002   0.26   0.76
2   0.90009   1.1   0.0001996404857   0.2601620081   0.759955
3   0.89991   1.1   0.0002003604863   0.2598380081   0.760045
...```

Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: no header

**Description**

See description of parent `custom.annotated`
7.2 method

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.49.41.1.3 interface_id

• Keywords Area
  • method
  • multifidelity_function_train
  • import_approx_points_file
  • custom_annotated
  • interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.49.41.2 annotated

• Keywords Area
  • method
  • multifidelity_function_train
  • import_approx_points_file
  • annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

  • file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn  nln_ineq_con_1 nln_ineq_con_2
1   NO_ID  0.9  1.1  0.0002  0.26  0.76
2   NO_ID  0.90009  1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID  0.89991  1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.49.41.3 freeform

- Keywords Area
- method
- multifidelity_function_train
- import_approx_points_file
- freeform

Selects freeform file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
...  0.9  1.1  0.0002  0.26  0.76
  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.49.41.4 active_only

- Keywords Area
Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.49.42 export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:
### Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

#### Usage Tips

Dakota exports tabular data in one of three formats:

- annotated *(default)*
- custom_annotated
- freeform

#### 7.2.49.42.1 custom_annotated

- Keywords Area
- method
- multifidelity_function_train
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

#### Topics

This keyword is related to the topics:

- file_formats

#### Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>Dakota Keyword</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>Dakota Keyword</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>Dakota Keyword</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1         0.9  1.1  0.0002  0.00026  0.76
2         0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3         0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2.49.42.1.1 header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.49.42.1.2 eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated
7.2.49.42.1.3 interface_id

- Keywords Area
- method
- multifidelity_function_train
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.49.42.2 annotated

- Keywords Area
- method
- multifidelity_function_train
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```environment
tabular_data
tabular_data_file = ’dakota_summary.dat’
annotated
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.00020003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.49.42.3 freeform

- Keywords Area

- method

- multifidelity_function_train

- export_approx_points_file

- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
 0.9  1.1  0.0002  0.26  0.76
 0.90009  1.1  0.0001996404857  0.2601620081  0.759955
 0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.49.43 seed_sequence

- Keywords Area
- method
  - multifidelity_function_train
- seed_sequence

Sequence of seed values for multi-stage random sampling
## 7.2 Specification

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** system-generated (non-repeatable)

### Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

### Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

### Usage Tips

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

### Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
    pilot_samples = 100 50 40 30 20
    seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

### See Also

These keywords may also be of interest:

- `seed`
- `fixed_seed`

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• method
• multifidelity_function_train
• fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

• block_pointer
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
        0.1 0.2 0.6
        0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
    id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
    id_method = 'DACE'
    model_pointer = 'DACE_M'
    sampling sample_type lhs
        samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
    id_interface = 'I1'
    system asynch evaluation_concurrency = 5
```
Keywords Area

analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.50 multilevel_function_train

- Keywords Area
- method
- multilevel_function_train

Multilevel uncertainty quantification using function train expansions

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td></td>
<td>Sample allocation approach for multilevel expansions</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>metric_scale</td>
<td></td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td>discrepancy_-emulation</td>
<td></td>
<td>Formulation for emulation of model discrepancies</td>
</tr>
<tr>
<td>Optional</td>
<td>Parameter</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>---------------------------</td>
<td>-------------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rounding_tolerance</td>
<td>An accuracy tolerance that is used to guide rounding during rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>arithmetic_-tolerance</td>
<td>A secondary rounding tolerance used for post-processing</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>regression_type</td>
<td>Type of solver for forming function train approximations by regression</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_-iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_cross_-iterations</td>
<td>Maximum number of iterations for cross-approximation during a rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>solver_tolerance</td>
<td>Convergence tolerance for the optimizer used during the regression solve.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>collocation_points-sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-----------------------------</td>
<td>---------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>start_order-sequence</td>
<td>Sequence of start orders used in a multi-stage expansion</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>adapt_order</td>
<td>Activate adaptive procedure for determining the best basis order</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>kick_order</td>
<td>Increment used when adapting the basis order in function train methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_order</td>
<td>Maximum polynomial order of each univariate function within the functional tensor train.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_cv_order-candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>start_rank-sequence</td>
<td>Sequence of start ranks used in a multi-stage expansion</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>adapt_rank</td>
<td>Activate adaptive procedure for determining best rank representation</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>kick_rank</td>
<td>The increment in rank employed during each iteration of the rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_rank</td>
<td>Limits the maximum rank that is explored during a rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_cv_rank_-candidates</td>
<td>Limit the number of cross-validation candidates for rank</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_-emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_-refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
<tr>
<td>-----------------------</td>
<td>-------------------------------------------------------------------------------------</td>
<td>---------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_-levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_-decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
<td></td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_-points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_-points_file</td>
<td>Output file for surrogate model value evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td>Sequence of seed values for multi-stage random sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
<td></td>
</tr>
</tbody>
</table>
As described in the function_train method and the function_train model, the function train (FT) approximation is a polynomial expansion that exploits low rank structure within the mapping from input random variables to output quantities of interest (QoI). For multilevel and multifidelity function train approximations, we decompose this expansion into several constituent expansions, one per model form or solution control level, where independent function train approximations are constructed for the low-fidelity/coarse resolution model and one or more levels of model discrepancy.

In a three-model case with low-fidelity (L), medium-fidelity (M), and high-fidelity (H) models and an additive discrepancy approach, we can denote this as:

\[ Q_H \approx \tilde{Q}^L + \tilde{\Delta}^{ML} + \tilde{\Delta}^{HM} \]

where \( \Delta^{ij} \) represents a discrepancy expansion computed from \( Q^i - Q^j \) and reduced rank representations of these discrepancies may be targeted (\( r_{HM} < r_{ML} < r_L \)).

In multilevel approaches, sample allocation for the constituent expansions is performed as described in allocation_control.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments (expansion moments only)
- Probability Density
- Level Mappings

In addition, the execution group has the attribute equiv_hf_evals, which records the equivalent number of high-fidelity evaluations.

**See Also**

These keywords may also be of interest:

- function_train
- function_train

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- max_function_evaluations

Sample allocation approach for multilevel expansions

Specification

Alias: none

Argument(s): none

Child Keywords:
Description

Multilevel expansions, including regression-based polynomial chaos expansion (PCE) and function train (FT) expansions, require a sample allocation strategy. Three options are currently available:

- allocation based on assuming a convergence rate for the estimator variance (for regression PCE)
- restricted isometry property (RIP) sampling (for regression PCE via compressed sensing)
- rank sampling (for FT)

Default Behavior

Current defaults, when allocation_control is not specified, are estimator variance for PCE and rank sampling for FT.

7.2.50.2.1 estimator_variance

- Keywords Area
- method
- multilevel_function_train
- allocation_control
- estimator_variance

Variance of mean estimator within multilevel polynomial chaos

Specification

Alias: none

Argument(s): none

Child Keywords:
Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

$$Var[\hat{Q}] = \frac{\sigma_Q^2}{N}$$

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

$$Var[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}$$

for free parameters $\gamma$ and $\kappa$, with default values that may be overridden as part of this specification block.

This approach is supported for regression-based PCE approaches (over-determined least squares, under-determined compressed sensing, or orthogonal least interpolation).

In practice, it can be challenging to estimate a smooth convergence rate for estimator variance in the presence of abrupt transitions in the quality of sparse recoveries. As a result, sample allocation by greedy refinement is generally preferred.

Examples

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the convergence of the estimator variance, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_polynomial_chaos
    orthogonal_matching_pursuit
    expansion_order_sequence = 2
    pilot_samples = 10
    collocation_ratio = .9
    allocation_control
      estimator_variance estimator_rate = 2.5
    seed = 1237
    convergence_tolerance = .01
```

7.2.50.2.1.1 estimator_rate

- Keywords Area
- method
- multilevel_function_train
- allocation_control
- estimator_variance
- estimator_rate

Rate of convergence of mean estimator within multilevel polynomial chaos
### 7.2 method

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 2

**Description**

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{\gamma^\kappa N}
\]

for free parameters \(\gamma\) and \(\kappa\).

The default values are \(\gamma = 1\) and \(\kappa = 2\) (adopts a more aggressive sample profile by assuming a faster convergence rate than Monte Carlo). This advanced specification option allows to user to specify \(\kappa\), overriding the default.

#### 7.2.50.2.2 rank_sampling

- **Keywords** Area  
- **method**  
- **multilevel_function_train**  
- **allocation_control**  
- **rank_sampling**

Sample allocation based on rank sampling within multilevel function train

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Multilevel function train based on regression may allocate the number of samples per level based on the collocation ratio times the regression size. The regression size is determined by the rank per core and the basis order per dimension as described at function_train, where these ranks and orders may be either user-specified values (for initial sample allocation), incremented values (for external adaptation by Dakota), or recovered values (in the case of internal C3 adaptation using adapt_rank).

The adaptive algorithm starts from a pilot sample, shapes the profile based on the regression size computed from the current orders and recovered ranks, and iterates until convergence.

This capability is ** experimental** and under active development.
Examples

This example starts with rank-two order-two initial expansion for each level, with twice as many samples as regression coefficients. As the recovered rank is updated for each level, as dictated by the internal adapt_rank approach, the number of samples is incremented as needed in order to synchronize with the specified collocation ratio. In this case, the basis order is fixed and only the ranks and associated samples are updated for each level.

```plaintext
method,
  model_pointer = 'HIERARCH'
  multifidelity_function_train
    allocation_control rank_sampling
      p_refinement uniform
        start_rank_sequence = 2 2 2 2
        adapt_rank   kick_rank = 2   max_rank = 10
        start_order_sequence = 2 2 2 2
        collocation_ratio = 2.  seed = 160415
        convergence_tolerance = 1.e-2
        max_refinement_iterations = 5
```

7.2.5.0.3 convergence_tolerance

- Keywords Area
- method
- multilevel_function_train
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:
- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPLLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.50.4 metric_scale

- Keywords Area
- method
- multilevel_function_train
- metric_scale

define scaling of statistical metrics when adapting UQ surrogates

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** relative

**Child Keywords:**

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<th>Description of Group scale of refinement metric (Group 1)</th>
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<th>Dakota Keyword Description</th>
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<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
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<tr>
<td></td>
<td>absolute</td>
<td></td>
<td>use of absolute statistical metrics when adapting UQ surrogates</td>
</tr>
</tbody>
</table>

**Description**

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The **absolute** option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The **relative** option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

The default is **relative**.

**Expected Output**
Usage Tips
Additional Discussion

7.2.50.4.1 relative

- Keywords Area
- method
- multilevel_function_train
- metric_scale
- relative

relative scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.50.4.2 absolute

- Keywords Area
- method
- multilevel_function_train
- metric_scale
- absolute

use of absolute statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none
Description

When adapting a UQ surrogate model, the absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.50.5 discrepancy_emulation

- Keywords Area
- method
- multilevel_function_train
- discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none

Argument(s): none

Default: distinct

Child Keywords:

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<tr>
<td>recursive</td>
<td>recursive</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
<td></td>
</tr>
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</table>

Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. distinct emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. recursive emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting
for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

Examples

```plaintext
distinct
```

Distinct formulation for emulation of model discrepancies.

Specification

**Alias:** paired

**Argument(s):** none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.50.5.2 recursive

Recursive formulation for emulation of model discrepancies.
The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.50.6 rounding_tolerance

- Keywords Area
- method
- multilevel_function_train
- rounding_tolerance

An accuracy tolerance that is used to guide rounding during rank adaptation.

Specification

Alias: none

Argument(s): none

Description

The rounding tolerance is used to measure how closely one can approximate a functional tensor train representation at a step of the optimization procedure using one of lower ranks. If this tolerance is small, then less compression is achieved but greater accuracy is obtained. If this tolerance is big, greater compression is achieved at the expense of accuracy.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- adapt_rank
- arithmetic_tolerance
7.2.50.7  arithmetic_tolerance


• Keywords Area

• method

• multilevel_function_train

• arithmetic_tolerance

A secondary rounding tolerance used for post-processing

Specification

Alias: none

Argument(s): REAL

Default: 1.e-10

Description

After forming one or more function train approximations, this tolerance is used for rolling up final results, in particular, combining multilevel/multifidelity expansions and forming products for higher-order moment estimation. It is generally intended as a looser tolerance than rounding_tolerance.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

• rounding_tolerance

7.2.50.8  regression_type


• Keywords Area

• method

• multilevel_function_train

• regression_type

Type of solver for forming function train approximations by regression

Specification

Alias: none

Argument(s): none

Child Keywords:
## Description

Function train approximations are formed based on regression for the set of coefficients described at `function_train`. Solver options include least squares and regularized regression, where the latter penalizes high-order terms to mitigate overfitting.

### Default Behavior

The default regression solver is least squares.

7.2.50.8.1 `ls`

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **regression_type**
- **ls**

Use least squares solver for forming function train approximations by regression

### Specification

**Alias**: none

**Argument(s)**: none

### Description

Least squares is the standard solver for forming function train approximations based on regression. This option does not include regularization.

7.2.50.8.2 `rls2`

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **regression_type**
- **rls2**

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<td>Use least squares solver for forming function train approximations by regression</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>rls2</code></td>
<td>Use regularized regression solver for forming function train approximations</td>
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</table>
Use regularized regression solver for forming function train approximations

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
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<td>Required</td>
<td></td>
<td>l2_penalty</td>
<td>Penalty value applied in regularized regression solver for function train approximations</td>
</tr>
</tbody>
</table>

**Description**

RLS2 is a regularized least squares solver that applies an L2 penalty to mitigate overfitting. It includes the option to specify a penalty parameter.

**See Also**

These keywords may also be of interest:

- l2_penalty

7.2.50.8.2.1 l2_penalty

- Keywords Area
- method
- multilevel_function_train
- regression_type
- rls2
- l2_penalty

Penalty value applied in regularized regression solver for function train approximations

**Specification**

**Alias:** none

**Argument(s):** REAL

**Description**

Regularized regression applies a penalty to aspects of the recovered solution (e.g., cardinality, higher-order oscillatory terms, et al.) to mitigate overfitting. This specification provides the scalar penalty parameter (its value is not currently adapted or cross-validated).
Default Behavior
If this penalty parameter is not specified, then we adopt the internal C3 default.

See Also
These keywords may also be of interest:

• rls2

7.2.50.9 max_solver_iterations

- Keywords Area
- method
- multilevel_function_train
- max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification
Alias: none
Argument(s): INTEGER
Default: 100

Description
When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.50.10 max_cross_iterations

- Keywords Area
- method
- multilevel_function_train
- max_cross_iterations

Maximum number of iterations for cross-approximation during a rank adaptation.

Specification
Alias: none
Argument(s): INTEGER
Default: 1
Description

The number of iterations for the cross-approximation algorithm is limited within the `adapt_rank` procedure.

See Also

These keywords may also be of interest:

- `adapt_rank`

7.2.50.11 `solver_tolerance`

- Keywords Area
- method
- multilevel_function_train
- `solver_tolerance`

Convergence tolerance for the optimizer used during the regression solve.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.0e-10

Description

Defines the function value and function gradient tolerance for optimizer used solving the regression problem for the functional tensor train coefficients.

7.2.50.12 `response_scaling`

- Keywords Area
- method
- multilevel_function_train
- `response_scaling`

Perform bounds-scaling on response values prior to surrogate emulation

Specification

**Alias:** none

**Argument(s):** none

**Default:** use original data
Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to \([0,1]\) based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.50.13 tensor_grid

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **tensor_grid**

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** regression with random sample set

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.50.14 collocation_points_sequence

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **collocation_points_sequence**

Sequence of collocation point counts used in a multi-stage expansion
Specification

Alias: pilot_samples

Argument(s): INTEGERLIST

Description

Each level entry of the \texttt{collocation_points_sequence} applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., \texttt{collocation_points}

See Also

These keywords may also be of interest:

- \texttt{collocation_points}

7.2.50.15 collocation_ratio

- Keywords Area
- method
- multilevel\_function\_train
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none

Argument(s): REAL

Default: 0.

Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the \texttt{collocation_ratio} allows for specification of a constant factor applied to \( N \) (e.g., \texttt{collocation\_ratio} = 2, produces samples = 2\( N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using \texttt{ratio\_order}. In this case, the number of samples becomes \( cN^o \) where \( c \) is the \texttt{collocation\_ratio} and \( o \) is the \texttt{ratio\_order}. The \texttt{use\_derivatives} flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o + 1 \)).
7.2 method  

7.2.50.16 start_order_sequence

- Keywords Area
- method
- multilevel_function_train
- start_order_sequence

Sequence of start orders used in a multi-stage expansion

Specification

**Alias:** order_sequence

**Argument(s):** INTEGERLIST

**Default:** 2

**Child Keywords:**

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<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
</tbody>
</table>

Description

Each level entry of the `start_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support start order sequences include multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `start_order`

See Also

These keywords may also be of interest:

- `start_order`

7.2.50.16.1 dimension_preference

- Keywords Area
- method
- multilevel_function_train
- start_order_sequence
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.50.17 adapt_order

- Keywords Area
- method
- multilevel_function_train
- adapt_order

Activate adaptive procedure for determining the best basis order

---

Specification

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

This option activates a cross validation procedure to select the best basis order for representing the current set of data. It searches from `start_order` to `max_order` in increments of `kick_order`.

**Default Behavior**

No cross validation for basis order.

**Examples**

This example shows specification of an order adaptation starting at order 2, incrementing by 2, and limited at order 6.
model,
    id_model = 'FT'
surrogate global function_train
start_rank = 4
    adapt_order start_order = 2 kick_order = 2 max_order = 6
solver_tolerance = 1e-12
rounding_tolerance = 1e-12
dace_method_pointer = 'SAMPLING'

Note that adapt_rank and adapt_order can either be combined or used separately.

See Also

These keywords may also be of interest:

* start_order
* kick_order
* max_order

### 7.2.50.18 kick_order

- Keywords Area
- method
- multilevel_function_train
- kick_order

increment used when adapting the basis order in function train methods

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1

**Description**

Used in combination with adapt_order, kick_order defines the size of the increment that is used during the cross validation process, within the range of start_order and max_order.

**Default Behavior**

Default is 1.

**See Also**

These keywords may also be of interest:

* adapt_order
7.2.50.19 max_order

- Keywords Area
- method
- multilevel_function_train
- max_order

Maximum polynomial order of each univariate function within the functional tensor train.

Specification

Alias: none
Argument(s): INTEGER
Default: unsigned short max

Description

This specification is inactive for current adaptation options. In future adaptation options (i.e., cross-approximation adaptation), this will specify the maximum order that can be obtained during adaptation, where polynomial order will start from start_order and be limited by max_order.

Default Behavior
Max order is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- start_order

7.2.50.20 max_cv_order_candidates

- Keywords Area
- method
- multilevel_function_train
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushorts max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment \_\_max\_order
- increment \_\_max\_rank\_order
- polynomial\_chaos
- multifidelity\_polynomial\_chaos
- function\_train
- multifidelity\_function\_train

7.2.50.21 start\_rank\_sequence

Sequence of start ranks used in a multi-stage expansion

Specification

Alias: rank\_sequence

Argument(s): INTEGERLIST

Default: 2

Description

Each level entry of the start\_rank\_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support start rank sequences include multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., start\_rank

See Also

These keywords may also be of interest:

- start\_rank
7.2.50.22  adapt_rank

- Keywords Area
- method
- multilevel_function_train
- adapt_rank

Activate adaptive procedure for determining best rank representation

Specification

**Alias:** none

**Argument(s):** none

**Default:** false

Description

The adaptive algorithm proceeds as follows:

1. Start from rank \texttt{start_rank} and form an approximation
2. Adapt the current approximation by searching for a solution with lower rank that achieves L2 accuracy within \texttt{epsilon} tolerance of the reference.
3. If a lower rank solution is found with comparable accuracy, then stop. If not, increase the rank by an amount specified by \texttt{kick_rank}.
4. Return to step 2 and continue until either \texttt{max_rank} is reached or a converged rank (rank less than current reference with comparable accuracy) is found.

Default Behavior

No cross validation for rank.

Examples

This example shows specification of a rank adaptation starting at rank 2, incrementing by 2, and limited at rank 10.

```dakota
model,
  id_model = 'FT'
  surrogate global function_train
  start_order = 5
  adapt_rank start_rank = 2  kick_rank = 2  max_rank = 10
  solver_tolerance = 1e-12
  rounding_tolerance = 1e-12
  dace_method_pointer = 'SAMPLING'
```

Note that \texttt{adapt_rank} and \texttt{adapt_order} can either be combined or used separately.

See Also

These keywords may also be of interest:

- \texttt{start_rank}
- \texttt{kick_rank}
- \texttt{max_rank}
7.2.50.23  kick_rank

- Keywords Area
- method
- multilevel_function_train
- kick_rank

The increment in rank employed during each iteration of the rank adaptation.

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The adapt_rank procedure increments the rank on each iteration, using an increment defined by kick_rank.

Default Behavior

Default is 1.

See Also

These keywords may also be of interest:

- adapt_rank

7.2.50.24  max_rank

- Keywords Area
- method
- multilevel_function_train
- max_rank

Limits the maximum rank that is explored during a rank adaptation.

Specification

Alias: none
Argument(s): INTEGER
Default: size_t max
The `adapt_rank` procedure increments the rank on each iteration, and the maximum value is limited by `max_rank`.

**Default Behavior**

Max rank is unbounded (adaptive sweep will stop when CV error increases).

**See Also**

These keywords may also be of interest:

- `adapt_rank`

7.2.50.25  `max_cv_rank_candidates`

- `Keywords Area`
- `method`
- `multilevel_function_train`
- `max_cv_rank_candidates`

Limit the number of cross-validation candidates for rank.

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Default**: `size_t max`

**Description**

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

**See Also**

These keywords may also be of interest:

- `increment_max_rank`
- `increment_max_rank_order`
- `function_train`
- `multifidelity_function_train`
7.2.50.26 samples_on_emulator

- Keywords Area
- method
- multilevel_function_train
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples
Argument(s): INTEGER

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.50.27 sample_type

- Keywords Area
- method
- multilevel_function_train
- sample_type

Selection of sampling strategy

Specification

Alias: none
Argument(s): none
Default: random
Child Keywords:
### Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

### Default Behavior

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

### Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

#### 7.2.50.27.1 lhs

- Keywords Area
- method
- multilevel_function_train
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

### Specification

**Alias:** none

**Argument(s):** none

### Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

### Default Behavior

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>
Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the \texttt{lhs} keyword must appear in conjunction with the \texttt{sample_type} keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```plaintext
method sampling
   sample_type lhs
   samples = 20
```

7.2.50.27.2 random

- Keywords Area
- method
- multilevel_function_train
- sample_type
- random

Uses purely random Monte Carlo sampling to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The \texttt{random} keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the \texttt{random} keyword must be specified in conjunction with the \texttt{sample_type} keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```plaintext
method sampling
   sample_type random
   samples = 200
```
7.2.50.28  rng

- Keywords Area
- method
- multilevel_function_train
- rng

Selection of a random number generator

Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.50.28.1  mt19937

- Keywords Area
- method
Generates random numbers using the Mersenne twister

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on Wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

**Default Behavior**

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

**Examples**

```plaintext
method
    sampling
        sample_type lhs
        samples = 10
        seed = 98765
        rng mt19937
```

Generates pseudo-random numbers using the Pecos package

**Specification**

**Alias**: none  
**Argument(s)**: none
**Description**

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

**7.2.50.29 probability_refinement**

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **probability_refinement**

Allow refinement of probability and generalized reliability results using importance sampling.

**Topics**

This keyword is related to the topics:

- **reliability_methods**

**Specification**

**Alias:** sample_refinement

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Importance Sampling Approach (Group 1)</td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>
## Description

The `probability_refinement` allows refinement of probability and generalized reliability results using importance sampling. If one specifies `probability_refinement`, there are some additional options. One can specify which type of importance sampling to use (`import`, `adapt_import`, or `mm_adapt_import`). Additionally, one can specify the number of refinement samples to use with `refinement_samples` and the seed to use with `seed`.

The `probability_refinement` density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

### 7.2.50.29.1 import

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **probability_refinement**
- **import**

Importance sampling option for probability refinement

### Specification

**Alias:** none

**Argument(s):** none

### Description

**import** centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

### 7.2.50.29.2 adapt_import

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **probability_refinement**
• **adapt_import**

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

*adapt_import* centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

7.2.50.29.3 **mm_adapt_import**

• **Keywords Area**

• **method**

• **multilevel_function_train**

• **probability_refinement**

• **mm_adapt_import**

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

*mm_adapt_import* starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then *mm_adapt_import* proceeds similarly to *adapt_import* (sample until convergence).

7.2.50.29.4 **refinement_samples**

• **Keywords Area**

• **method**

• **multilevel_function_train**

• **probability_refinement**

• **refinement_samples**

Number of samples used to refine a probability estimate or sampling design.
7.2 Method

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial samples in a sampling design.

7.2.50.30 final_moments

- Keywords Area
- method
- multilevel_function_train
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>
Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```

7.2.50.30.1 none

- Keywords Area
- method
- multilevel_function_train
- final_moments
7.2 method

- none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method, sampling
   samples = 50 seed = 1234
   response_levels = 3.6e+11 1.2e+05 3.5e+05
   final_moments none

7.2.50.30.2 standard

- Keywords Area
- method
- multilevel_function_train
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.
Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, 
sampling 
  samples = 50  seed = 1234 
  response_levels = 3.6e+11 1.2e+05 3.5e+05 
  final_moments central 
```

**Keywords Area**

• method  
• multilevel_function_train  
• final_moments  
• central

Values at which to estimate desired statistics for each response
### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

### Description

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

### Default Behavior

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

### Expected Outputs

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

### Usage Tips

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

### Examples

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num-`
Keywords Area

_**response_levels**_ key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method,
    sampling,
        samples = 100 seed = 1
    complementary distribution
        response_levels = 3.6e+11 4.0e+11 4.4e+11
                        6.0e+04 6.3e+04 7.0e+04
                        3.5e+05 4.0e+05 4.5e+05
variables,
    normal_uncertain = 2
        means = 248.89, 593.33
        std_deviations = 12.4, 29.7
        descriptors = 'TF1n', 'TF2n'
    uniform_uncertain = 2
        lower_bounds = 199.3, 474.63
        upper_bounds = 298.5, 712.
        descriptors = 'TF1u', 'TF2u'
    weibull_uncertain = 2
        alphas = 12., 30.
        betas = 250., 590.
        descriptors = 'TF1w', 'TF2w'
    histogram_bin_uncertain = 2
        num_pairs = 3 4
        abscissas = 5 8 10 .1 .2 .3 .4
        counts = 17 21 0 12 24 12 0
        descriptors = 'TF1h', 'TF2h'
    histogram_point_uncertain
        real = 1
        num_pairs = 2
        abscissas = 3 4
        counts = 1 1
        descriptors = 'TF3h'
interface,
    system async evaluation_concurrency = 5
    analysis_driver = 'text_book'
responses,
    response_functions = 3
        no_gradients
        no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

```plaintext
Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>
PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>
PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844608686e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
</tbody>
</table>
```

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7.2 method

4.0000000000e+05  4.5000000000e+05  1.8000000000e-06

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
Response Level  Probability Level  Reliability Index  General Rel Index
--------------  -----------------  -----------------  -----------------
3.6000000000e+11  5.5000000000e-01
4.0000000000e+11  3.8000000000e-01
4.4000000000e+11  2.3000000000e-01

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
Response Level  Probability Level  Reliability Index  General Rel Index
--------------  -----------------  -----------------  -----------------
6.0000000000e+04  6.1000000000e-01
6.5000000000e+04  2.9000000000e-01
7.0000000000e+04  9.0000000000e-02

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
Response Level  Probability Level  Reliability Index  General Rel Index
--------------  -----------------  -----------------  -----------------
3.5000000000e+05  5.2000000000e-01
4.0000000000e+05  9.0000000000e-02
4.5000000000e+05  0.0000000000e+00

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.50.31.1  num_response_levels

- Keywords Area
- method
- multilevel_function_train
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.
Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.50.31.2  compute

- Keywords Area
- method
- multilevel_function_train
- response_levels
- compute

Selection of statistics to compute at each response level

Specification

Alias: none
Argument(s): none
Default: probabilities
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>probabilities</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>reliabilities</td>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>gen_reliabilities</td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
</tbody>
</table>
### Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

### Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

### Expected Output

The type of statistics specified by `compute` will be reported for each response level.

### Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

### Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

### Specification

**Alias:** none

**Argument(s):** none

Optional system reliability (series or parallel)
Description

The **probabilities** keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If **response_levels** is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the **probabilities** keyword should be specified in conjunction with the **compute** keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

### 7.2.50.31.2.2 reliabilities

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **response_levels**
- **compute**
- **reliabilities**

Computes reliabilities associated with response levels

**Specification**

**Alias:** none

**Argument(s):** none

Description

The **reliabilities** keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If **response_levels** is specified, the reliabilities are not computed by default. To change this behavior, the **reliabilities** keyword should be specified in conjunction with the **compute** keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
Examples

```dakota
method
    sampling
        sample_type random
        samples = 100 seed = 1
    complementary distribution
        response_levels = 3.6e+11 4.0e+11 4.4e+11
        6.0e+04 6.5e+04 7.0e+04
        3.5e+05 4.0e+05 4.5e+05
    compute gen_reliabilities
```

7.2.50.31.2.3 gen_reliabilities

- Keywords Area
- method
- multilevel_function_train
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

**Alias:** none

**Argument(s):** none

Description

The *gen_reliabilities* keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If *response_levels* is specified, the generalized reliabilities are not computed by default. To change this behavior, the *gen_reliabilities* keyword should be specified in conjunction with the *compute* keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```dakota
method
    sampling
        sample_type random
        samples = 100 seed = 1
    complementary distribution
        response_levels = 3.6e+11 4.0e+11 4.4e+11
        6.0e+04 6.5e+04 7.0e+04
        3.5e+05 4.0e+05 4.5e+05
    compute gen_reliabilities
```
7.2.50.31.2.4  system

- Keywords Area
- method
- multilevel_function_train
- response_levels
- compute
- system

Compute system reliability (series or parallel)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

**Description**

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.50.31.2.5  series

- Keywords Area
- method
- multilevel_function_train
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system
7.2 method

Specification

Alias: none
Argument(s): none

Description

See parent keyword `system` for description.

7.2.50.31.2.6 parallel

- Keywords Area
- method
- multilevel_function_train
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword `system` for description.

7.2.50.32 probability_levels

- Keywords Area
- method
- multilevel_function_train
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:
### Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

### Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

### Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```plaintext
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
  1. .8 .5 0.
  1. .3 .2 0.
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = ’TF1n’ ’TF2n’
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = ’TF1u’ ’TF2u’
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = ’TF1w’ ’TF2w’
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = ’TF1h’ ’TF2h’
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = ’TF3h’
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = ’text_book’
responses,
  response_functions = 3
```

### Keywords Area

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.1454122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431514744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028365236e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600556346e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td>6.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td>3.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431514744e+04</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td>8.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td>5.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td>3.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td>2.0000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.50.32.1 num_probability_levels

- Keywords Area
- method
• multilevel_function_train
• probability_levels
• num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.50.33 reliability_levels

• Keywords Area
• method
• multilevel_function_train
• reliability_levels

Specify reliability levels at which the response values will be estimated

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.50.33.1 num_reliability_levels

- Keywords Area
- method
- multilevel_function_train
- reliability_levels
- num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.50.34 gen_reliability_levels

- Keywords Area
- method
- multilevel_function_train
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:
### Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

### Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

#### 7.2.50.34.1 num_gen_reliability_levels

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **gen_reliability_levels**
- **num_gen_reliability_levels**

Specify which `gen_reliability_levels` correspond to which response

### Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** `gen_reliability_levels` evenly distributed among response functions

### Description

See parent page

#### 7.2.50.35 distribution

- **Keywords Area**
- **method**
7.2 method

- multilevel_function_train
- distribution

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.50.35.1 cumulative

- Keywords Area
- method
- multilevel_function_train
Keywords Area

- **distribution**
- **cumulative**

Computes statistics according to cumulative functions

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the **cumulative** keyword must appear in conjunction with the **distribution** keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```
method
  sampling
    sample_type lh
    samples = 10
    distribution cumulative
```

7.2.50.35.2 **complementary**

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **distribution**
- **complementary**

Computes statistics according to complementary cumulative functions

**Specification**

**Alias**: none

**Argument(s)**: none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```plaintext
method	sampling
  sample_type lhs
  samples = 10
  distribution complementary
```

7.2.50.36 variance_based_decomp

- Keywords Area
- method
- multilevel_function_train
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

**Alias:** none

**Argument(s):** none

**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
</tbody>
</table>
Optional

<table>
<thead>
<tr>
<th>drop_tolerance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

**Default Behavior**

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

**Expected Outputs**

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

**Usage Tips**

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.

**Examples**

```plaintext
method,
  polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2
```

**Theory**

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76] : "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].

7.2.50.36.1 interaction_order

- Keywords Area
- method
- multilevel_function_train
7.2 method

- variance_based_decomp
- interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

**Description**

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.50.36.2 drop_tolerance

- Keywords Area
- method
- multilevel_function_train
- variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed

**Description**

The drop_tolerance keyword allows the user to specify a value below which sensitivity indices generated by variance_based_decomp are not displayed.

**Default Behavior**

By default, all sensitivity indices generated by variance_based_decomp are displayed.

**Usage Tips**

For polynomial_chaos, which outputs main, interaction, and total effects by default, the univariate_effects may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
Examples

```
method, sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
  drop_tolerance = 0.001
```

7.2.50.37 diagonal_covariance

- Keywords Area
- method
- multilevel_function_train
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

**Alias**: none

**Argument(s)**: none

**Default**: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.50.38 full_covariance

- Keywords Area
- method
- multilevel_function_train
- full_covariance

Display the full covariance matrix

Specification

**Alias**: none

**Argument(s)**: none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.
7.2.50.39 import_approx_points_file

• Keywords Area

• method

• multilevel_function_train

• import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td></td>
<td></td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td>freeform</td>
<td></td>
<td></td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

**Default Behavior** No import of points at which to evaluate the surrogate.

**Expected Output** The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_approx_points_file = 'import.mcmc_annot.dat'
    annotated
```

7.2.50.39.1 custom_annotated

• Keywords Area

• method

• multilevel_function_train

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and *eval_id* (no *interface_id*), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

```
%eval_id x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9   1.1   0.0002  0.26   0.76
2  0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3  0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...
```

**7.2.50.39.1.1 header**

• *Keywords Area*
  • *method*
  • *multilevel_function_train*
  • *import_approx_points_file*
  • *custom_annotated*
  • *header*

Enable header row in custom-annotated tabular file

**Specification**

Alias: none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent *custom_annotated*

**7.2.50.39.1.2 eval_id**

• *Keywords Area*
  • *method*
  • *multilevel_function_train*
  • *import_approx_points_file*
  • *custom_annotated*
  • *eval_id*

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent *custom_annotated*

7.2.50.39.1.3 interface_id

- Keywords Area
- method
- multilevel_function_train
- import_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent *custom_annotated*

7.2.50.39.2 annotated

- Keywords Area
- method
- multilevel_function_train
- import_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.50.39.3 freeform

- **Keywords Area**
  - method
  - multilevel_function_train
  - import_approx_points_file

Selects freeform file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.50.39.4 active_only

- Keywords Area
Import only active variables from tabular data file

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

**export_approx_points_file**

**Specification**

**Alias:** export_points_file

**Argument(s):** STRING

**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate's predicted approximation to the truth model responses at those points.

### Usage Tips

Dakota exports tabular data in one of three formats:

- `annotated` (default)
- `custom_annotated`
- `freeform`

### 7.2.50.40.1 `custom_annotated`

- **Keywords Area**
- **method**
- **multilevel_function_train**
- **export_approx_points_file**
- **custom_annotated**

Selects custom-annotated tabular file format

### Topics

This keyword is related to the topics:

- `file_formats`

### Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

evironment
   tabular_data
   tabular_data_file = 'dakota_summary.dat'
   custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
7.2.50.40.1.1 header

- Keywords Area
- method
- multilevel_function_train
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.50.40.1.2 eval_id

- Keywords Area
- method
- multilevel_function_train
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated
7.2.50.40.1.3 interface_id

- Keywords Area
- method
- multilevel_function_train
- export_approx_points_file
- custom.annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent custom.annotated

7.2.50.40.2 annotated

- Keywords Area
- method
- multilevel_function_train
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.50.40.3 freeform

- Keywords Area
- method
- multilevel_function_train
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
7.2 method

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.50.41 seed_sequence

• Keywords Area
• method
• multilevel_function_train
• seed_sequence

Sequence of seed values for multi-stage random sampling
Specification

Alias: none

Argument(s): INTEGERLIST

Default: system-generated (non-repeatable)

Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
   model_pointer = 'HIERARCH'
   multilevel_sampling
   pilot_samples = 100 50 40 30 20
   seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

See Also

These keywords may also be of interest:

- seed
- fixed_seed

7.2.50.42 fixed_seed

- Keywords Area
Reuses the same seed value for multiple random sampling sets

Specification

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run

**Description**

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

**Examples**

```plaintext
method
sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

**Topics**

This keyword is related to the topics:

- `block_pointer`
Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
7.2 method

    analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.51 polynomial_chaos

    • Keywords Area
    • method
    • polynomial_chaos

Uncertainty quantification using polynomial chaos expansions

Specification

Alias: nond_polynomial_chaos

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Chaos coefficient estimation approach (Group 1)</td>
<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>metric_scale</td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_-iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
<table>
<thead>
<tr>
<th>Optional (Choose One)</th>
<th>Basis Polynomial Family (Group 2)</th>
<th>askey</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
</tr>
</tbody>
</table>

- **sparse_grid_level**: Level to use in sparse grid integration or interpolation.
- **cubature_integrand**: Cubature using Stroud rules and their extensions.
- **expansion_order**: The (initial) order of a polynomial expansion.
- **orthogonal_least_interpolation**: Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.
- **import_expansion_file**: Build a Polynomial Chaos Expansion (PCE) by import coefficients and a multi-index from a file.

- **askey**: Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
- **wiener**: Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.
<table>
<thead>
<tr>
<th>Optional</th>
<th>normalized</th>
<th>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>export_expansion-_file</td>
<td>Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_--emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate).</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy.</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator.</td>
</tr>
<tr>
<td>Optional</td>
<td>probability_--refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling.</td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response.</td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value.</td>
</tr>
</tbody>
</table>
### Keywords Area

<table>
<thead>
<tr>
<th>Optional</th>
<th>reliability_levels</th>
<th>Specify reliability levels at which the response values will be estimated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 3)</td>
<td>diagonal_covariance</td>
</tr>
<tr>
<td>Optional</td>
<td>full_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate model value evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

### Description

The polynomial chaos expansion (PCE) is a general framework for the approximate representation of random response functions in terms of finite-dimensional series expansions in standardized random variables.
\[ R = \sum_{i=0}^{P} \alpha_i \Psi_i(\xi) \]

where \( \alpha_i \) is a deterministic coefficient, \( \Psi_i \) is a multidimensional orthogonal polynomial and \( \xi \) is a vector of standardized random variables. An important distinguishing feature of the methodology is that the functional relationship between random inputs and outputs is captured, not merely the output statistics as in the case of many nondeterministic methodologies.

**Basis polynomial family (Group 1)**

Group 1 keywords are used to select the type of basis, \( \Psi_i \), of the expansion. Three approaches may be employed:

- **Wiener**: employs standard normal random variables in a transformed probability space, corresponding to Hermite orthogonal basis polynomials (see `wiener`).
- **Askey**: employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space, corresponding to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal basis polynomials, respectively (see `askey`).
- **Extended (default if no option is selected)**: The Extended option avoids the use of any nonlinear variable transformations by augmenting the Askey approach with numerically-generated orthogonal polynomials for non-Askey probability density functions. Extended polynomial selections replace each of the sub-optimal Askey basis selections for bounded normal, lognormal, bounded lognormal, loguniform, triangular, gumbel, frechet, weibull, and bin-based histogram.

For supporting correlated random variables, certain fallbacks must be implemented.

- The Extended option is the default and supports only Gaussian correlations.
- If needed to support prescribed correlations (not under user control), the Extended and Askey options will fall back to the Wiener option on a per variable basis. If the prescribed correlations are also unsupported by Wiener expansions, then Dakota will exit with an error.

Refer to `variable_support` for additional information on supported variable types, with and without correlation.

**Coefficient estimation approach (Group 2)**

To obtain the coefficients \( \alpha_i \) of the expansion, seven options are provided:

1. multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with `quadrature_order` and, optionally, `dimension_preference`).
2. multidimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`).
3. multidimensional integration by Stroud cubature rules and extensions as specified with `cubature_integrand`.
4. multidimensional integration by Latin hypercube sampling (specified with `expansion_order` and `expansion_samples`).
5. linear regression (specified with `expansion_order` and either `collocation_points` or `collocation_ratio`), using either over-determined (least squares) or under-determined (compressed sensing) approaches.
6. orthogonal least interpolation (specified with `orthogonal_least_interpolation` and `collocation_points`).
7. coefficient import from a file (specified with `import_expansion_file`). The expansion can be comprised of a general set of expansion terms, as indicated by the multi-index annotation within the file.
It is important to note that, for polynomial chaos using a single model fidelity, `quadrature_order`, `sparse_grid_level`, and `expansion_order` are scalar inputs used for a single expansion estimation. These scalars can be augmented with a `dimension_preference` to support anisotropy across the random dimension set. This differs from the use of sequence arrays in advanced use cases such as multilevel and multifidelity polynomial chaos, where multiple grid resolutions can be employed across a model hierarchy.

**Active Variables**

The default behavior is to form expansions over aleatory uncertain continuous variables. To form expansions over a broader set of variables, one needs to specify `active` followed by `state`, `epistemic`, `design`, or `all` in the variables specification block.

For continuous design, continuous state, and continuous epistemic uncertain variables included in the expansion, Legendre chaos bases are used to model the bounded intervals for these variables. However, these variables are not assumed to have any particular probability distribution, only that they are independent variables. Moreover, when probability integrals are evaluated, only the aleatory random variable domain is integrated, leaving behind a polynomial relationship between the statistics and the remaining design/state/epistemic variables.

**Covariance type (Group 3)**

These two keywords are used to specify how this method computes, stores, and outputs the covariance of the responses. In particular, the diagonal covariance option is provided for reducing post-processing overhead and output volume in high dimensional applications.

**Optional Keywords regarding method outputs**

Each of these sampling specifications refer to sampling on the PCE approximation for the purposes of generating approximate statistics.

- `sample_type`
- `samples`
- `seed`
- `fixed_seed`
- `rng`
- `probability_refinement`
- `distribution`
- `reliability_levels`
- `response_levels`
- `probability_levels`
- `gen_reliability_levels`

which should be distinguished from simulation sampling for generating the PCE coefficients as described in options 4, 5, and 6 above (although these options will share the `sample_type`, `seed`, and `rng` settings, if provided).

When using the `probability_refinement` control, the number of refinement samples is not under the user’s control (these evaluations are approximation-based, so management of this expense is less critical). This option allows for refinement of probability and generalized reliability results using importance sampling.

**Usage Tips**

If \( n \) is small (e.g., two or three), then tensor-product Gaussian quadrature is quite effective and can be the preferred choice. For moderate to large \( n \) (e.g., five or more), tensor-product quadrature quickly becomes too expensive and the sparse grid and regression approaches are preferred. Random sampling for coefficient estimation is generally not recommended due to its slow convergence rate. For incremental studies, approaches 4 and 5 support reuse of previous samples through the `incremental_lhs` and `reuse_points` specifications, respectively.

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = \)
2l + 1 are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension- _adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

**Additional Resources**


**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments
- Probability Density
- Level Mappings
- Variance-Based Decomposition (Sobol' Indices)

**Examples**

```python
method, polynomial_chaos
    sparse_grid_level = 2
    samples = 10000 seed = 12347 rng rnum2
    response_levels = .1 1. 50. 100. 500. 1000.
    variance_based_decomp
```

**See Also**

These keywords may also be of interest:

- stoch_collocation
- function_train

**7.2.51.1 p_refinement**

- Keywords Area
- method
- polynomial_chaos
- p_refinement

Automatic polynomial order refinement
Specification

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>p-refinement Type (Group 1)</td>
<td>uniform</td>
<td>refine an expansion uniformly in all dimensions.</td>
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<tr>
<td></td>
<td></td>
<td>dimension_adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
</tbody>
</table>

**Description**

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the `generalized` `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
7.2 method

7.2.51.1 uniform

- Keywords Area
- method
- polynomial_chaos
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.51.2 dimension_adaptive

- Keywords Area
- method
- polynomial_chaos
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

#### 7.2.51.1.2.1 sobol

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

---

#### Specification

**Alias**: none

**Argument(s)**: none

**Default**: generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.51.1.2.2 decay

- Keywords Area
- method
- polynomial_chaos
- p_refinement
- dimension_adaptive
- decay

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

Specification

Alias: none
Argument(s): none

Description

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.

Examples

```
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement decay
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.51.1.2.3 generalized

- Keywords Area
- method
- polynomial_chaos
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method,  
  polynomial_chaos  
  sparse_grid_level = 3  
  dimension_adaptive p_refinement generalized  
  max_iterations = 20  
  convergence_tol = 1.e-4
```

7.2.51.2 **max_refinement_iterations**

**Keywords Area**

- **method**
- **polynomial_chaos**
- **max_refinement_iterations**

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

method, polynomial_chaos
dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1

7.2.51.3 convergence_tolerance

- Keywords Area
- method
- polynomial_chaos
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.51.4 metric_scale

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **metric_scale**

define scaling of statistical metrics when adapting UQ surrogates

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** relative

### Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group scale of refinement metric (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>chose One</td>
<td></td>
<td>relative</td>
<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>absolute</td>
<td></td>
<td>absolute</td>
<td>use of absolute statistical metrics when adapting UQ surrogates</td>
</tr>
</tbody>
</table>

#### Description

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The `absolute` option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The `relative` option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

#### Default Behavior

The default is `relative`.

#### Expected Output

#### Usage Tips

#### Additional Discussion

### 7.2.51.4.1 relative

- **Keywords Area**
- **method**
relative scaling of statistical metrics when adapting UQ surrogates

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When adapting a UQ surrogate model, the `relative` option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**

---

use of absolute statistical metrics when adapting UQ surrogates

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When adapting a UQ surrogate model, the `absolute` option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**
7.2.51.5 quadrature_order

- Keywords Area
- method
- polynomial_chaos
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
</tbody>
</table>

*Optional/Choose One*

| Optional | Quadrature Rule Nesting (Group 1) | nested | non_nested | Enforce use of nested quadrature rules if available |

**Description**

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order, and, optionally, dimension_preference). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a dimension_preference, where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified quadrature_order and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n}(p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the quadrature_order specification supports an array input.
A corresponding sequence specification is documented at, e.g., quadrature_order_sequence and quadrature_order_sequence

See Also

These keywords may also be of interest:

- quadrature_order_sequence and method-multifidelity_stoch_collocation-quadrature_order_sequence

7.2.51.5.1 dimension_preference

- Keywords Area
- method
- polynomial_chaos
- quadrature_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none
Argument(s): REALLIST
Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.51.5.2 nested

- Keywords Area
- method
- polynomial_chaos
- quadrature_order
- nested

Enforce use of nested quadrature rules if available
Specification

Alias: none
Argument(s): none
Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.51.5.3 non_nested
  • Keywords Area
  • method
  • polynomial_chaos
  • quadrature_order
  • non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.51.6 sparse_grid_level
  • Keywords Area
  • method
  • polynomial_chaos
  • sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

Alias: none
Argument(s): INTEGER
Child Keywords:
### Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User's Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User's Manual for additional details). As for `quadrature_order`, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`.

### See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
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<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 1)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>unrestricted</td>
<td>Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
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</table>

---
7.2.51.6.1  dimension_preference

- Keywords Area
- method
- polynomial_chaos
- sparse_grid_level
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none

Argument(s): REALLIST

Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.51.6.2  restricted

- Keywords Area
- method
- polynomial_chaos
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

Alias: none

Argument(s): none

Default: restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.51.6.3 unrestricted

- **Keywords** Area
- method
- polynomial_chaos
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.51.6.4 nested

- **Keywords** Area
- method
- polynomial_chaos
- sparse_grid_level
Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.51.6.5 non_nested

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **sparse_grid_level**
- **non_nested**

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.51.7 cubature_integrand

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **cubature_integrand**

Cubature using Stroud rules and their extensions

**Specification**

**Alias:** none  
**Argument(s):** INTEGER
Description

Multi-dimensional integration by Stroud cubature rules [80] and extensions [94], as specified with `cubature_integrand`. A total-order expansion is used, where the isotropic order \( p \) of the expansion is half of the integrand order, rounded down. The total number of terms \( N \) for an isotropic total-order expansion of order \( p \) over \( n \) variables is given by

\[
N = 1 + P = 1 + \sum_{s=1}^{p} \frac{1}{s!} \prod_{r=0}^{s-1} (n+r) = \frac{(n+p)!}{n!p!}
\]

Since the maximum integrand order is currently five for normal and uniform and two for all other types, at most second- and first-order expansions, respectively, will be used. As a result, cubature is primarily useful for global sensitivity analysis, where the Sobol’ indices will provide main effects and, at most, two-way interactions. In addition, the random variable set must be independent and identically distributed (\( iid \)), so the use of `askey` or `wiener` transformations may be required to create \( iid \) variable sets in the transformed space (as well as to allow usage of the higher order cubature rules for normal and uniform). Note that global sensitivity analysis often assumes uniform bounded regions, rather than precise probability distributions, so the \( iid \) restriction would not be problematic in that case.

7.2.51.8 expansion_order

- Keywords Area
- method
- polynomial_chaos
- expansion_order

The (initial) order of a polynomial expansion

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
| Optional          | dimension_-
preference        |                | A set of weights specifying the relative importance of each uncertain variable (dimension) |
| Optional          | basis_type           |                | Specify the type of basis truncation to be used for a Polynomial Chaos Expansion. |
### Description

When the expansion order for a polynomial chaos expansion is specified, the coefficients may be computed by integration based on random samples or by regression using either random or sub-sampled tensor product quadrature points.

Multidimensional integration by Latin hypercube sampling (specified with `expansion_samples`). In this case, the expansion order \( p \) cannot be inferred from the numerical integration specification and it is necessary to provide an `expansion_order` to specify \( p \) for a total-order expansion.

Linear regression (specified with either `collocation_points` or `collocation_ratio`). A total-order expansion is used and must be specified using `expansion_order` as described in the previous option. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the `collocation_ratio` allows for specification of a constant factor applied to \( N \) (e.g., `collocation_ratio = 2.` produces samples = \( 2N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using `ratio_order`. In this case, the number of samples becomes \( cN^\alpha \) where \( c \) is the `collocation_ratio` and \( \alpha \) is the `ratio_order`. The `use_derivatives` flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^\alpha + 1 \)). When admissible, a constrained least squares approach is employed in which response values are first reproduced exactly and error in reproducing response derivatives is minimized. Two collocation grid options are supported: the default is Latin hypercube sampling ("point collocation"), and an alternate approach of "probabilistic collocation" is also available through inclusion of the `tensor_grid` keyword. In this alternate case, the collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

If `collocation_points` or `collocation_ratio` is specified, the PCE coefficients will be determined by regression. If no regression specification is provided, appropriate defaults are defined. Specifically SVD-based least-squares will be used for solving over-determined systems and under-determined systems will be solved using LASSO. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using the least squares method, incorporating the constraint that the PCE coefficients match the function values at the collocation points. The number of samples required becomes \( cN^\alpha + 1 \) when admissible.

### Table

<table>
<thead>
<tr>
<th>Required (Choose One)</th>
<th>Group 1</th>
<th>collocation_points</th>
<th>Number of collocation points used to estimate expansion coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_samples</td>
<td>Number of simulation samples used to estimate the expected value of a set of PCE coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
system will be solved using equality constrained least squares. Technical information on the various methods listed below can be found in the Linear regression section of the Theory Manual. Some of the regression methods (O-MP, LASSO, and LARS) are able to produce a set of possible PCE coefficient vectors (see the Linear regression section in the Theory Manual). If cross validation is inactive, then only one solution, consistent with the noise_tolerance, will be returned. If cross validation is active, Dakota will choose between possible coefficient vectors found internally by the regression method across the set of expansion orders (1,...,expansion_order) and the set of specified noise tolerances and return the one with the lowest cross validation error indicator.

7.2.51.8.1  dimension_preference

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none
Argument(s): REALLIST
Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.51.8.2  basis_type

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.
Specification

Alias: none

Argument(s): none

Child Keywords:
### 7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td><strong>PCE Basis Type (Group 1)</strong></td>
<td><strong>tensor_product</strong></td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>total_order</strong></td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>adapted</strong></td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

### Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.

#### 7.2.51.8.2.1 tensor_product

- **Keywords Area**
  - method
  - polynomial_chaos
  - expansion_order
  - basis_type
  - tensor_product

Use a tensor-product index set to construct a polynomial chaos expansion.

#### Specification

- **Alias**: none
- **Argument(s)**: none

#### Description

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$
Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Dakota Keyword</th>
<th>Description Description</th>
</tr>
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<tbody>
<tr>
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<td></td>
</tr>
<tr>
<td>Optional</td>
<td>advancements</td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>----------</td>
<td>--------------</td>
<td>--------------------------------------------------------</td>
</tr>
<tr>
<td>Optional</td>
<td>soft_convergence_limit</td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

**7.2.51.8.2.4 advancements**

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

### Specification

**Alias**: none

**Argument(s)**: INTEGER

### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

**7.2.51.8.2.5 soft_convergence_limit**

- Keywords Area
- method
- polynomial_chaos
The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

**7.2.51.8.3 collocation_points**

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points

Number of collocation points used to estimate expansion coefficients

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Child Keywords:**

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<thead>
<tr>
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<th>Description of Group</th>
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<tr>
<td><strong>Optional (Choose One)</strong></td>
<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td>Method</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>least_angle_regression</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>cross_validation</td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>ratio_order</td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
</tr>
<tr>
<td>Keywords Area</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>-------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.

A corresponding sequence specification is documented at, e.g., `collocation_points_sequence` and `collocation_points_sequence`.

**See Also**

These keywords may also be of interest:

- `collocation_points_sequence` method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence`

7.2.51.8.3.1 least_squares

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_squares

Compute the coefficients of a polynomial expansion using least squares.
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>LSQ Regression Approach (Group 1)</td>
<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>equality_-constrained</td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.51.8.3.2 svd

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **collocation_points**
- **least_squares**
- **svd**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
7.2.51.8.3.3 equality_constrained

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_squares
- equality_constrained

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.51.8.3.4 orthogonal_matching_pursuit

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Requires/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.51.8.3.5 noise_tolerance

- Keywords Area
- method
- polynomial_choas
- expansion_order
- collocation_points
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none
Argument(s): REALLIST
Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.3.6 basis_pursuit

- Keywords Area
- method
- polynomial_choas
- expansion_order
- collocation_points
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

Specification

Alias: bp
Argument(s): none
### Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

#### 7.2.51.8.3.7 basis_pursuit_denoising

- **Keywords Area**
  - method
  - polynomial_chaos
  - expansion_order
  - collocation_points
  - basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

### Specification

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

### Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

#### 7.2.51.8.3.8 noise_tolerance

- **Keywords Area**
  - method
  - polynomial_chaos
  - expansion_order
  - collocation_points
  - basis_pursuit_denoising
  - noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
7.2 method

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.3.9 least_angle_regression

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.51.8.3.10 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_angle_regression
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.3.11 least_absolute_shrinkage

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso  
**Argument(s):** none  

**Child Keywords:**

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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td>The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.
7.2.51.8.3.12 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias**: none

**Argument(s)**: REALIST

**Default**: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.3.13 l2_penalty

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias**: none

**Argument(s)**: REAL

**Default**: 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
7.2.51.8.3.14  cross_validation

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- cross_validation

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_-candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

Description

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.51.8.3.15  noise_only

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.
7.2 method

Specification

Alias: none
Argument(s): none
Default: false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.51.8.3.16 max_cv_order_candidates

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_points
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
7.2.51.8.3.17 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none

Argument(s): REAL

Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^\text{ratio_order}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.

7.2.51.8.3.18 response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

Specification

Alias: none

Argument(s): none

Default: use original data
Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

### 7.2.51.8.3.19 use_derivatives

- **Keywords Area**
- `method`
- `polynomial_chaos`
- `expansion_order`
- `collocation_points`
- `use_derivatives`

Use derivative data to construct surrogate models

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

### 7.2.51.8.3.20 tensor_grid

- **Keywords Area**
- `method`
- `polynomial_chaos`
- `expansion_order`
- `collocation_points`
• **tensor_grid**

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.51.8.3.21 **reuse_points**

• **Keywords Area**
• **method**
• **polynomial_chaos**
• **expansion_order**
• **collocation_points**
• **reuse_points**

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

7.2.51.8.3.22 **max_solver_iterations**

• **Keywords Area**
• **method**
• **polynomial_chaos**
• **expansion_order**
• collocation_points
• max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none

Argument(s): INTEGER

Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.51.8.4 collocation_ratio

• Keywords Area
  • method
  • polynomial_chaos
  • expansion_order
  • collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none

Argument(s): REAL

Child Keywords:
| Basis Pursuit | Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming. |
| Basis Pursuit Denoising | Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization. |
| Least Angle Regression | Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method. |
| Least Absolute Shrinkage | Compute the coefficients of a polynomial expansion by using the LASSO problem. |
| Optional | Cross Validation | Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion. |
| Optional | Ratio Order | Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients. |
### 7.2 method

<table>
<thead>
<tr>
<th>Optional</th>
<th>response_scaling</th>
<th>Perform bounds-scaling on response values prior to surrogate emulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
</tr>
</tbody>
</table>

**Description**

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o \)).

**7.2.51.8.4.1 least_squares**

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **collocation_ratio**
- **least_squares**

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias**: none
**Argument(s):** none  
**Default:** svd

### Child Keywords:

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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>equality_-constrained</td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

### Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.51.8.4.2 svd

- Keywords Area
- method
- polynomial_choas
- expansion_order
- collocation_ratio
- least_squares
- svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

### Specification

**Alias:** none  
**Argument(s):** none

### Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
7.2.51.8.4.3 equality_constrained

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- least_squares
- equality_constrained

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Specification

Alias: none

Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.51.8.4.4 orthogonal_matching_pursuit

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.51.8.4.5 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none
Argument(s): REALLIST
Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.4.6 basis_pursuit

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

Specification

Alias: bp
Argument(s): none
Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.51.8.4.7 basis_pursuit_denoising

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

Specification

Alias: bpdn

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
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<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.51.8.4.8 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.4.9 least_angle_regression

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

**Alias:** lars

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.51.8.4.10 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- least_angle_regression
• noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.4.11 least_absolute_shrinkage

• Keywords Area
• method
• polynomial_chaos
• expansion_order
• collocation_ratio
• least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso

Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td>The l2 penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.
7.2.51.8.4.12 noise_tolerance

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.51.8.4.13 l2_penalty

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

Description

The $l_2$ penalty used when performing compressed sensing with elastic net.
7.2.51.8.4.14  cross_validation

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- cross_validation

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_only</td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_cv_order_candidates</td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.51.8.4.15  noise_only

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.
Specification

**Alias:** none

**Argument(s):** none

**Default:** false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.51.8.4.16  max_cv_order_candidates

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
• multifidelity_polynomial_chaos
• function_train
• multifidelity_function_train

7.2.51.8.4.17 ratio_order

• Keywords Area
• method
• polynomial_chaos
• expansion_order
• collocation_ratio
• ratio_order

Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none

Argument(s): REAL

Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.

7.2.51.8.4.18 response_scaling

• Keywords Area
• method
• polynomial_chaos
• expansion_order
• collocation_ratio
• response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

Specification

Alias: none

Argument(s): none

Default: use original data
Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- multifidelity_polynomial_chaos
- multifidelity_function_train

7.2.51.8.4.19 use_derivatives

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.51.8.4.20 tensor_grid

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- collocation_ratio
7.2 method

- **tensor_grid**

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.51.8.4.21 reuse_points

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **collocation_ratio**
- **reuse_points**

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

7.2.51.8.4.22 max_solver_iterations

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
• collocation_ratio
• max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none

Argument(s): INTEGER

Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.51.8.5 expansion_samples

• Keywords Area
• method
• polynomial_chaos
• expansion_order
• expansion_samples

Number of simulation samples used to estimate the expected value of a set of PCE coefficients

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

Description

In this case, the expansion order $p$ cannot be inferred from the numerical integration specification and it is necessary to provide an `expansion_order` to specify $p$ for a total-order expansion.

A corresponding sequence specification is documented at, e.g., `expansion_samples_sequence` and `expansion_sequence`
See Also

These keywords may also be of interest:

- `expansion_samples_sequence`
- `method-multilevel_polynomial_chaos`
- `expansion_order_sequence`
- `expansion_samples_sequence`

7.2.51.8.5.1 reuse_points

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **expansion_samples**
- **reuse_points**

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.51.8.6 import_build_points_file

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```python
    method
        polynomial_chaos
            expansion_order = 4
            import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.51.8.6.1 custom_annotated

- Keywords Area
  - method
  - polynomial_chaos
  - expansion_order
  - import_build_points_file
  - custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether `header`, `eval_id`, and `interface_id` column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
**Keywords Area**

**environment**

**tabular_data**

*tabular_data_file = 'dakota_summary.dat'

*custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.10009</td>
<td>0.0002</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.10009</td>
<td>0.0002</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### 7.2.51.8.6.2 header

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **import_build_points_file**
- **custom_annotated**
- **header**

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent `custom_annotated`

### 7.2.51.8.6.3 eval_id

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **expansion_order**
- **import_build_points_file**
- **custom_annotated**
- **eval_id**

Enable evaluation ID column in custom-annotated tabular file
7.2 method

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.51.8.6.4 interface_id

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.51.8.6.5 annotated

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- import_build_points_file
- annotated

Selects annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
7.2.51.8.6.6 freeform

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.51.6.7 active_only

- Keywords Area
- method
- polynomial_chaos
- expansion_order
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.51.9 orthogonal_least_interpolation

- Keywords Area
- method
• polynomial_chaos
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>collocation_points</td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods expansion_order cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

7.2.51.9.1 collocation_points

• Keywords Area
• method
• polynomial_chaos
• orthogonal_least_interpolation
• collocation_points

Number of collocation points used to estimate expansion coefficients
Specification

Alias: none
Argument(s): INTEGER

Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches. A corresponding sequence specification is documented at, e.g., collocation_points_sequence and collocation_points_sequence.

See Also

These keywords may also be of interest:

- collocation_points_sequence
- method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence

7.2.51.9.2 tensor_grid

Keywords Area

- method
- polynomial_chaos
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): INTEGERLIST
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.51.9.3 reuse_points

Keywords Area

- method
- polynomial_chaos
- orthogonal_least_interpolation
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
### 7.2 method

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The *reuse_points* option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify *reuse_points* so that any points that have been previously generated (for example, from the *import_points_file*) can be reused.

#### 7.2.51.9.4 import_build_points_file

- **Keywords Area**
  - method
  - polynomial_chaos
  - orthogonal_least_interpolation
  - import_build_points_file

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The *import_build_points_file* allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g.
surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```plaintext
method
  polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.51.9.4.1 custom_annotated

- Keywords Area
- method
- polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

### Table

<table>
<thead>
<tr>
<th>Optional</th>
<th></th>
<th>Optional</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>header</td>
<td></td>
<td>eval_id</td>
<td></td>
</tr>
<tr>
<td>interface_id</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>7.2.51.9.4.2</th>
<th>header</th>
</tr>
</thead>
<tbody>
<tr>
<td>Keywords Area</td>
<td></td>
</tr>
<tr>
<td>method</td>
<td></td>
</tr>
</tbody>
</table>
Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no header

**Description**

See description of parent `custom_annotated`

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`
7.2.51.9.4.4  interface_id

- Keywords Area
- method
- polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

 Alias: none
 Argument(s): none
 Default: no interface_id column

Description

See description of parent custom_annotated

7.2.51.9.4.5  annotated

- Keywords Area
- method
- polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

 Alias: none
 Argument(s): none
 Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  NO_ID  0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.51.9.4.6 freeform

- Keywords Area
- method
- polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
        freeform
```

Resulting tabular file:

```
  0.9     1.1      0.0002    0.26     0.76
  0.90009 1.1 0.0001996404857 0.2601620081 0.759955
  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
  ...
```

Import only active variables from tabular data file

- Keywords Area
- method
- polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- active_only
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.51.10 import_expansion_file

- Keywords Area
- method
- polynomial_chaos
- import_expansion_file

Build a Polynomial Chaos Expansion (PCE) by import coefficients and a multi-index from a file

Specification

Alias: none

Argument(s): STRING

Description

The coefficients can be specified in an arbitrary order. The multi-index provided is used to generate a sparse expansion that consists only of the indices corresponding to the non-zero coefficients provided in the file.

7.2.51.11 askey

- Keywords Area
- method
- polynomial_chaos
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- `polynomial_chaos`
- `askey`

### 7.2.51.13 normalized

- `Keywords Area`
- `method`
- `polynomial_chaos`
- `normalized`

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

**Description**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

### 7.2.51.14 export_expansion_file

- `Keywords Area`
- `method`
- `polynomial_chaos`
- `export_expansion_file`

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.

**Specification**

**Alias:** none

**Argument(s):** STRING

**Description**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.
7.2.51.15  samples_on_emulator

- Keywords Area
- method
- polynomial_chaos
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias:  samples
Argument(s):  INTEGER

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
  polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.51.16  sample_type

- Keywords Area
- method
- polynomial_chaos
- sample_type

Selection of sampling strategy

Specification

Alias:  none
Argument(s):  none
Default:  random
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```method
sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

**7.2.51.16.1 lhs**

- Keywords Area
- method
- polynomial_chaos
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**
Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

Usage Tips
Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples
```
method sampling
   sample_type lhs
   samples = 20
```

7.2.51.16.2 random
- Keywords Area
- method
- polynomial_chaos
- sample_type
- random

Uses purely random Monte Carlo sampling to sample variables

Specification
- **Alias**: none
- **Argument(s)**: none

Description
The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior
In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

Usage Tips
Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples
```
method sampling
   sample_type random
   samples = 200
```
Selection of a random number generator

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** Mersenne twister (mt19937)

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```
method
    sampling
        sample_type lhs
        samples = 10
        seed = 98765
        rng rnum2
```

7.2.51.17.1 mt19937

- Keywords Area
- method
Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

**Default Behavior**

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

Generates pseudo-random numbers using the Pecos package

**Specification**

**Alias:** none  
**Argument(s):** none
Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.51.18 probability_refinement

- Keywords Area
- method
- polynomial_chaos
- probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: `sample_refinement`

Argument(s): none

Default: no refinement

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Importance Sampling Approach (Group 1)</td>
<td><code>import</code></td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>
### Description

The `probability_refinement` allows refinement of probability and generalized reliability results using importance sampling. If one specifies `probability_refinement`, there are some additional options. One can specify which type of importance sampling to use (`import`, `adapt_import`, or `mm_adapt_import`). Additionally, one can specify the number of refinement samples to use with `refinement_samples` and the seed to use with `seed`.

The `probability_refinement` density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

#### 7.2.51.18.1 import

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **probability_refinement**
- **import**

Importance sampling option for probability refinement

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

`import` centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

#### 7.2.51.18.2 adapt_import

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **probability_refinement**
• **adapt_import**

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

**7.2.51.18.3 mm_adapt_import**

• **Keywords Area**
  • method
  • polynomial_ chaos
  • probability_refinement
  • mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).

**7.2.51.18.4 refinement_samples**

• **Keywords Area**
  • method
  • polynomial_ chaos
  • probability_refinement
  • refinement_samples

Number of samples used to refine a probability estimate or sampling design.
7.2 method

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using uni-modal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.

7.2.51.19 final_moments

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **final_moments**

Output moments of the specified type and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

**Default:** standard

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td><strong>none</strong></td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>standard</strong></td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>
Output central moments and include them within the set of final statistics.

**Description**

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
        secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
    nested
        sub_method_pointer = 'UQ'
        primary_response_mapping = 1. 1. 0.
        secondary_response_mapping = 0. 0. 1.
```

7.2.51.19.1 none

- Keywords Area
- method
- polynomial_chaos
- final_moments
7.2 method

- none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method, sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none

7.2.51.19.2 standard

- Keywords Area
- method
- polynomial_chaos
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.
Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

Values at which to estimate desired statistics for each response
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_response_-levels</td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>compute</td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num-
The Dakota input file below specifies a sampling method with response levels of interest.

```
method,
  sampling,
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    ... 6.0e+04 6.5e+04 7.0e+04
                       3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
    num_pairs = 3 4
    abscissas = .5, 8, 10, .1, .2, .3, .4
    counts = 17 21 0 12 24 12 0
    descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

```
Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844608686e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
</tbody>
</table>
```
7.2 method

4.0000000000e+05  4.5000000000e+05  1.8000000000e-06

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.51.20.1  num_response_levels

- Keywords Area
- method
- polynomial_chaos
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.
Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```plaintext
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.251.20.2 compute

- Keywords Area
- method
- polynomial_chaos
- response_levels
- compute

Selection of statistics to compute at each response level

Specification

Alias: none
Argument(s): none
Default: probabilities

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
<td></td>
</tr>
<tr>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
<td></td>
</tr>
<tr>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
<td></td>
</tr>
</tbody>
</table>
Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```plaintext
method sampling
    sample_type random
    samples = 100 seed = 1
complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
        6.0e+04 6.5e+04 7.0e+04
        3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```

7.2.51.20.2.1 probabilities

- Keywords Area
- method
- polynomial_chaos
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

Specification

**Alias**: none

**Argument(s)**: none
Description

The probabilities keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the probabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

7.2.51.20.2.2 reliabilities

- Keywords Area

Specification

Alias: none

Argument(s): none

Description

The reliabilities keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the reliabilities are not computed by default. To change this behavior, the reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
Examples

```plaintext
method
sampling
  sample_type random
  samples = 100 seed = 1
complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
compute reliabilities
```

7.2.51.20.2.3 gen_reliabilities

- Keywords Area
- method
- polynomial_chaos
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

**Alias:** none

**Argument(s):** none

Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
sampling
  sample_type random
  samples = 100 seed = 1
complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
compute gen_reliabilities
```
Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Choose One)</td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td>parallel</td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.51.20.2.5 series

- Keywords Area
- method
- polynomial_chaos
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system
7.2 method

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.51.20.2.6 parallel

• Keywords Area
• method
• polynomial_chaos
• response_levels
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.51.21 probability_levels

• Keywords Area
• method
• polynomial_chaos
• probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute
Child Keywords:
Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
    1. .8 .5 0.
    1. .3 .2 0.
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond to the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.51.21.1 num_probability_levels

- Keywords Area
- method
• polynomial_chaos
• probability_levels
• num_probability_levels

Specify which probability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** probability_levels evenly distributed among response functions

**Description**

See parent page

7.2.51.22 reliability_levels

  • Keywords Area
  • method
  • polynomial_chaos
  • reliability_levels

Specify reliability levels at which the response values will be estimated

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.51.22.1 num_reliability_levels

- Keywords Area
- method
- polynomial_chaos
- reliability_levels
- num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none
Argument(s): INTEGERLIST
Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.51.23 gen_reliability_levels

- Keywords Area
- method
- polynomial_chaos
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute
Child Keywords:
Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.51.23.1 num_gen_reliability_levels

- Keywords Area
- method
- polynomial_chaos
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.51.24 distribution

- Keywords Area
- method
• polynomial_chaos

• distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative

7.2.51.24.1 cumulative

• Keywords Area

• method

• polynomial_chaos
Computes statistics according to cumulative functions

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the **cumulative** keyword must appear in conjunction with the **distribution** keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

---

**Complementary Area**

• **Keywords Area**  
• **method**  
• **polynomial_choas**  
• **distribution**  
• **complementary**

Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none  
**Argument(s):** none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

7.2.51.25  variance_based_decomp

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **variance_based_decomp**

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

**Alias:** none

**Argument(s):** none

**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
</tbody>
</table>
Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

Default Behavior

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol’ indices.

Expected Outputs

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.

Examples

```
method, polynomial_chaos # or stoch_collocation
sparse_grid_level = 3
variance_based_decomp interaction_order = 2
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76] : “The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input.”

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].

7.2.51.25.1 interaction_order

- Keywords Area
- method
- polynomial_chaos
• variance_based_decomp
• interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

Specification

Alias: none
Argument(s): INTEGER
Default: Unrestricted (VBD includes all interaction orders present in the expansion)

Description

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification).

Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.51.25.2 drop_tolerance

• Keywords Area
• method
• polynomial_chaos
• variance_based_decomp
• drop_tolerance

 Suppresses output of sensitivity indices with values lower than this tolerance

Specification

Alias: none
Argument(s): REAL
Default: All VBD indices displayed

Description

The drop_tolerance keyword allows the user to specify a value below which sensitivity indices generated by variance_based_decomp are not displayed.

Default Behavior

By default, all sensitivity indices generated by variance_based_decomp are displayed.

Usage Tips

For polynomial_chaos, which outputs main, interaction, and total effects by default, the univariate_effects may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
Examples

method,
sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
drop_tolerance = 0.001

7.2.51.26 diagonal_covariance

- Keywords Area
- method
- polynomial_chaos
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none

Argument(s): none

Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.51.27 full_covariance

- Keywords Area
- method
- polynomial_chaos
- full_covariance

Display the full covariance matrix

Specification

Alias: none

Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.
import_approx_points_file

- Keywords Area
- method
- polynomial_chaos
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

**Specification**

**Alias**: none

**Argument(s)**: STRING

**Default**: no point import from a file

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

**Default Behavior** No import of points at which to evaluate the surrogate.

**Expected Output** The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```plaintext
method
  polynomial_chaos
    expansion_order = 4
    import_approx_points_file = 'import.mcmc_annot.dat'
    annotated
```

```
7.2.51.28 custom_annotated

- Keywords Area
- method
- polynomial_chaos
```
• `import_approx_points_file`

• `custom_annotated`

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• `file_formats`

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

```eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.51.28.1.1 header

• Keywords Area

• method

• polynomial_chaos

• import_approx_points_file

• custom_annotated

• header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated

7.2.51.28.1.2 eval_id

• Keywords Area

• method

• polynomial_chaos

• import_approx_points_file

• custom_annotated

• eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

*See description of parent* custom_annotated

7.2.51.28.1.3  interface_id

- Keywords Area
- method
- polynomial_chaos
- import_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

*See description of parent* custom_annotated

7.2.51.28.2  annotated

- Keywords Area
- method
- polynomial_chaos
- import_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

evironment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90099 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...

7.2.51.28.3 freeform

• Keywords Area
  • method
  • polynomial_chaos
  • import_approx_points_file
  • freeform

Selects freeform file format
This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.51.28.4 active_only

- Keywords Area
7.2 method

- method
- polynomial_chaos
- import_approx_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.51.29 export_approx_points_file

- Keywords Area
- method
- polynomial_chaos
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

### Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

#### 7.2.51.29.1 custom_annotated

- Keywords Area
- method
- polynomial_chaos
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

### Topics

This keyword is related to the topics:

- file_formats

### Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**
7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.9009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
7.2.51.29.1.1  header

- Keywords Area
- method
- polynomial_chaos
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no header

**Description**

See description of parent custom_annotated

7.2.51.29.1.2  eval_id

- Keywords Area
- method
- polynomial_chaos
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent custom_annotated
7.2.51.29.1.3 interface_id

- Keywords Area
- method
- polynomial_choas
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no interface_id column

Description

See description of parent custom_annotated

7.2.51.29.2 annotated

- Keywords Area
- method
- polynomial_choas
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
evironment tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.51.29.3 freeform

- Keywords Area
- method
- polynomial_chaos
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
7.2 method

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```dakota
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.51.30 seed

• Keywords Area
  • method
  • polynomial_chaos
  • seed

Seed of the random number generator
Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.51.31 fixed_seed

- Keywords Area
- method
- polynomial_chaos
- fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The
same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

**7.2.51.32 model_pointer**

- **Keywords Area**
- **method**
- **polynomial_chaos**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.
Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
  0.1 0.2 0.6
  0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.52 multifidelity_polynomial_chaos

• Keywords Area
• method
  • multifidelity_polynomial_chaos

Multifidelity uncertainty quantification using polynomial chaos expansions

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_-iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>metric_scale</td>
<td>Define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>statistics_mode</td>
<td>Type of statistical metric roll-up for multifidelity UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_-emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Chaos Coefficient Estimation Approach (Group 1)</td>
<td>quadrature_order_-sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level_-sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td>Expansion Order Sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------------------------</td>
<td>-------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Orthogonal Least Interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Basis Polynomial Family (Group 2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
<td></td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>Normalized</td>
<td>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>Export Expansion File</td>
<td>Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</td>
<td></td>
</tr>
<tr>
<td>Parameter</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------</td>
<td>-------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>samples_on_emulator</strong></td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>sample_type</strong></td>
<td>Selection of sampling strategy</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>rng</strong></td>
<td>Selection of a random number generator</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>probability_refinement</strong></td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>final_moments</strong></td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>response_levels</strong></td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>probability_levels</strong></td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>reliability_levels</strong></td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>gen_reliability_levels</strong></td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>distribution</strong></td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| Optional | **Covariance Type** (Group 3) | Optional/(Choose One) | Optional
---|---|---|---
**variance_based_decomp** | diagonal_covariance | import_approx_points_file | export_approx_points_file
**full_covariance** | | seed_sequence | fixed_seed
**model_pointer** | | | |

### Description

As described in *polynomial_chaos*, the polynomial chaos expansion (PCE) is a general framework for the approximate representation of random response functions in terms of series expansions in standardized random variables:

\[
R = \sum_{i=0}^{P} \alpha_i \Psi_i(\xi)
\]

where \( \alpha_i \) is a deterministic coefficient, \( \Psi_i \) is a multidimensional orthogonal polynomial and \( \xi \) is a vector of standardized random variables.

In the multilevel and multifidelity cases, we decompose this expansion into several constituent expansions, one per model form or solution control level. In a bi-fidelity case with low-fidelity (LF) and high-fidelity (HF) models and an additive discrepancy approach, we have:

\[
R = \sum_{i=0}^{P_{LF}} \alpha_i^{LF} \Psi_i(\xi) + \sum_{i=0}^{P_{HF}} \delta \Psi_i(\xi)
\]

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
where $\delta_i$ is a coefficient for the discrepancy expansion.

The same specification options are available as described in polynomial_chaos with one key difference: many of the coefficient estimation inputs change from a scalar input for a single expansion to a sequence specification for a low-fidelity expansion followed by multiple discrepancy expansions.

To obtain the coefficients $\alpha_i$ and $\delta_i$ for each of the expansions, the following options are provided:

1. multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with `quadrature_order_sequence`, and, optionally, `dimension_preference`).
2. multidimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level_sequence` and, optionally, `dimension_preference`).
3. multidimensional integration by Latin hypercube sampling (specified with `expansion_order_sequence` and `expansion_samples_sequence`).
4. linear regression (specified with `expansion_order_sequence` and either `collocation_points_sequence` or `collocation_ratio`), using either over-determined (least squares) or under-determined (compressed sensing) approaches.
5. orthogonal least interpolation (specified with `orthogonal_least_interpolation` and `collocation_points_sequence`).

It is important to note that, while `quadrature_order_sequence`, `sparse_grid_level_sequence`, `expansion_order_sequence`, `expansion_samples_sequence`, and `collocation_points_sequence` are array inputs, only one scalar from these arrays is active at a time for a particular expansion estimation. In order to specify anisotropy in resolution across the random variable set, a `dimension_preference` specification can be used to augment scalar specifications for quadrature order, sparse grid level, and expansion order.

Multifidelity UQ using PCE requires that the model selected for iteration by the method specification is a multifidelity surrogate model (see hierarchical), which defines an `ordered_model_sequence` (see hierarchical). Two types of hierarchies are supported: (i) a hierarchy of model forms composed from more than one model within the `ordered_model_sequence`, or (ii) a hierarchy of discretization levels comprised from a single model within the `ordered_model_sequence` which in turn specifies a `solution_level_control` (see `solution_level_control`).

In both cases, an expansion will first be formed for the low fidelity model or coarse discretization, using the first value within the coefficient estimation sequence, along with any specified refinement strategy. Second, expansions are formed for one or more model discrepancies (the difference between response results if additive correction or the ratio of results if multiplicative correction), using all subsequent values in the coefficient estimation sequence (if the sequence does not provide a new value, then the previous value is reused) along with any specified refinement strategy. The number of discrepancy expansions is determined by the number of model forms or discretization levels in the hierarchy.

After formation and refinement of the constituent expansions, each of the expansions is combined (added or multiplied) into an expansion that approximates the high fidelity model, from which the final set of statistics are generated. For polynomial chaos expansions, this high fidelity expansion can differ significantly in form from the low fidelity and discrepancy expansions, particularly in the multiplicative case where it is expanded to include all of the basis products.

Additional Resources

Dakota provides access to multifidelity PCE methods through the NonDMultilevelPolynomialChaos class. Refer to the Stochastic Expansion Methods chapter of the Theory Manual[16] for additional information on the Multifidelity PCE algorithm.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments (expansion moments only)
In addition, the execution group has the attribute `equiv_hf_evals`, which records the equivalent number of high-fidelity evaluations.

**Examples**

```dakota
method, multifidelity_polynomial_chaos  
   model_pointer = 'HIERARCH'  
   sparse_grid_level_sequence = 4 3 2

model,  
   id_model = 'HIERARCH'  
   surrogate hierarchical  
   ordered_model_fidelities = 'LF' 'MF' 'HF'  
   correction additive zeroth_order
```

**See Also**

These keywords may also be of interest:

- `polynomial_chaos`
- `multifidelity_stoch_collocation`
- `multifidelity_function_train`

7.2.52.1  **p_refinement**

- Keywords Area
- method
- multifidelity_polynomial_chaos
- p_refinement

Automatic polynomial order refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group p-refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>uniform</td>
<td>uniform</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
</tbody>
</table>
### 7.2.52.1 uniform

- **Keywords**
  - Area

- **method**

- **multifidelity_polynomial_chaos**

- **p_refinement**

- **uniform**

Refine an expansion uniformly in all dimensions.

<table>
<thead>
<tr>
<th>dimension_adaptive</th>
<th>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.</th>
</tr>
</thead>
</table>

**Description**

The **p_refinement** keyword specifies the usage of automated polynomial order refinement, which can be either **uniform** or **dimension_adaptive**.

The **dimension_adaptive** option is supported for the tensor-product quadrature and Smolyak sparse grid options and **uniform** is supported for tensor and sparse grids as well as regression approaches (**collocation_points** or **collocation_ratio**).

Each of these refinement cases makes use of the **max_iterations** and **convergence_tolerance** method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The **dimension_adaptive** case can be further specified to utilize **sobol**, **decay**, or **generalized** refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (**sobol** case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (**decay** case: low decay rates result in high dimension preference). In these two cases as well as the **uniform** refinement case, the **quadrature_order** or **sparse_grid_level** are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the **uniform** refinement case with regression approaches, the **expansion_order** is ramped by one on each iteration while the oversampling ratio (either defined by **collocation_ratio** or inferred from **collocation_points** based on the initial expansion) is held fixed. Finally, the **generalized** **dimension_adaptive** case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of **p_refinement** or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.52.1.2 dimension_adaptive

- Keywords Area
- method
- multifidelity_polynomial_chaos
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>Dimension Adaptivity Estimation Approach (Group 1)</td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.</td>
</tr>
</tbody>
</table>
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

#### 7.2.52.1.2.1 sobol

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** generalized

#### Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.

**Examples**

```dakota
method, polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```

7.2.52.1.2.3 **generalized**

- **Keywords Area**
  - method
  - multifidelity_polynomial_chaos
  - p_refinement
  - dimension_adaptive

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.
7.2 method

Specification

**Alias:** none

**Argument(s):** none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.52.2 max_refinement_iterations

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **max_refinement_iterations**

Maximum number of expansion refinement iterations

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

```
method, polynomial_chaos
  dimension_adaptive generalized
  p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```
7.2.52.3 convergence_tolerance

- Keywords Area
- method
- multifidelity_polynomial_chaos
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration.

For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.52.4 metric_scale

- Keywords Area
- method
• **multifidelity_polynomial_chaos**

• **metric_scale**

Define scaling of statistical metrics when adapting UQ surrogates

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** relative

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group scale of refinement metric (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td><strong>relative</strong></td>
<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td></td>
<td><strong>absolute</strong></td>
<td>use of absolute statistical metrics when adapting UQ surrogates</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The **absolute** option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The **relative** option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

The default is **relative**.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.52.4.1 relative

• **Keywords Area**

• **method**

• **multifidelity_polynomial_chaos**

• **metric_scale**

• **relative**

Relative scaling of statistical metrics when adapting UQ surrogates
Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.52.4.2 absolute

• Keywords Area
• method
• multifidelity_polynomial_chaos
• metric_scale
• absolute

use of absolute statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.52.5 statistics_mode

• Keywords Area
• method
• multifidelity_polynomial_chaos
• statistics_mode

type of statistical metric roll-up for multifidelity UQ methods
### Specification

**Alias:** none

**Argument(s):** none

**Default:** DEFAULT_EXPANSION_STATS

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>active</td>
<td>active</td>
<td>use statistics for the active expansion within multifidelity refinement</td>
</tr>
<tr>
<td></td>
<td>combined</td>
<td>combined</td>
<td>use statistics for the combined expansion within multifidelity refinement</td>
</tr>
</tbody>
</table>

### Description

When performing adaptive refinement in multifidelity UQ methods, one can adapt the expansion for a model level/fidelity based on its local statistics (active) or based on the statistics for the aggregate expansion (combined). The latter case may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

**Default Behavior**

Default for multifidelity methods is combined statistics.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.52.5.1 active

- Keywords Area
  - method
  - multifidelity_polynomial_chaos
  - statistics_mode
  - active

use statistics for the active expansion within multifidelity refinement

### Specification

**Alias:** none

**Argument(s):** none
Keywords Area

Description

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the active model level/fidelity. This approach avoids some computational overhead in some cases by avoiding expansion roll-up.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.52.5.2 combined

• Keywords Area

• method

• multifidelity_polynomial_chaos

• statistics_mode

• combined

use statistics for the combined expansion within multifidelity refinement

Specification

Alias: none

Argument(s): none

Description

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the aggregate multifidelity expansion. This approach may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.52.6 allocation_control

• Keywords Area

• method

• multifidelity_polynomial_chaos

• allocation_control

Sample allocation approach for multifidelity expansions
### Specification

**Alias:** none  
**Argument(s):** none

#### Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required</strong> (Choose One)</td>
<td>Multifidelity Sample Allocation Control (Group 1)</td>
<td>greedy</td>
<td>Sample allocation based on greedy refinement within multifidelity polynomial chaos</td>
</tr>
</tbody>
</table>

#### Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

#### Default Behavior

The default, when `allocation_control` is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

#### 7.2.52.6.1 greedy

- **Keywords Area**  
  - method  
  - `multifidelity_polynomial_chaos`  
  - `allocation_control`  
  - greedy

Sample allocation based on greedy refinement within multifidelity polynomial chaos

#### Specification

**Alias:** none  
**Argument(s):** none

#### Description

Multifidelity polynomial chaos supports greedy refinement strategies, spanning regression and projection approaches for computing expansion coefficients. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within a unified competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any $c/p/\beta/\beta^*$ level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.
Examples

The following example of greedy multifidelity regression starts from a zeroth-order reference expansion (a constant) for each level, and generates candidate refinements for each level that are competed in an integrated greedy competition. The number of new samples for the incremented candidate expansion order is determined from the collocation ratio. In this case, the number of candidates for each level is limited to one uniform refinement of the current expansion order.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
  allocation_control greedy
  p_refinement uniform
  expansion_order_sequence = 0
  collocation_ratio = .9 seed = 160415
  orthogonal_matching_pursuit
  convergence_tolerance 1.e-2
```

The next example employs generalized sparse grids within a greedy multifidelity competition. Each modeling level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all model levels is competed within an integrated greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each model level and the number of candidates grows rapidly with random dimension and grid level.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
  allocation_control greedy
  p_refinement dimension_adaptive generalized
  sparse_grid_level_sequence = 0 unrestricted
  convergence_tolerance 1.e-8
```

7.2.52.7 discrepancy_emulation

- Keywords Area
- method
- multifidelity_polynomial_chaos
- discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

- **Alias**: none
- **Argument(s)**: none
- **Default**: distinct
- **Child Keywords**:
Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. distinct emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. recursive emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

Examples

```
method,  multilevel_polynomial_chaos
         expansion_order_sequence = 2
         collocation_ratio = .9
         orthogonal_matching_pursuit
         discrepancy_emulation recursive
```

7.2.52.7.1 distinct

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **discrepancy_emulation**
- **distinct**

Distinct formulation for emulation of model discrepancies.

Specification

**Alias:** paired

**Argument(s):** none
Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.52.7.2 recursive

- Keywords Area
- method
- multifidelity_polynomial_chaos
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.52.8 quadrature_order_sequence

- Keywords Area
- method
- multifidelity_polynomial_chaos
- quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST
Child Keywords:
### Description

Each level entry of the `quadrature_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `quadrature_order`.

### See Also

These keywords may also be of interest:

- `quadrature_order`

### 7.2.52.8.1 dimension_preference

- **Keywords Area**
- **method**
- `multifidelity_polynomial_chaos`
- `quadrature_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

### Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.52.8.2 nested

- Keywords Area
- method
- multifidelity_polynomial_chaos
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none
Argument(s): none
Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.52.8.3 non_nested

- Keywords Area
- method
- multifidelity_polynomial_chaos
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule
7.2.52.9  sparse_grid_level_sequence

- Keywords Area
- method
- multifidelity_polynomial_chaos
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Optional</td>
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<td></td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 1)</td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
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<tr>
<td></td>
<td></td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>unrestricted</td>
<td>Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

**Description**

Each level entry of the *sparse_grid_level_sequence* applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., *sparse_grid_level*
See Also

These keywords may also be of interest:

- `sparse_grid_level`

7.2.52.9.1 `dimension_preference`

- Keywords Area
- method
- `multifidelity_polynomial_chaos`
- `sparse_grid_level_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- `sobol`
- `decay`

7.2.52.9.2 `restricted`

- Keywords Area
- method
- `multifidelity_polynomial_chaos`
- `sparse_grid_level_sequence`
- `restricted`

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid ( \( i = 4l + 1 \) ) or an order \( m \) for a tensor grid ( \( i = 2m - 1 \) ). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.52.9.3 unrestricted

- Keywords Area
- method
- multifidelity_polynomial_chaos
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid ( \( i = 4l + 1 \) ) or an order \( m \) for a tensor grid ( \( i = 2m - 1 \) ). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.52.9.4 nested

- Keywords Area
- method
- multifidelity_polynomial_chaos
- sparse_grid_level_sequence
• nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.52.9.5 non_nested

• Keywords Area

• method

• multifidelity_polynomial_chaos

• sparse_grid_level_sequence

• non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none

Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.52.10 expansion_order_sequence

• Keywords Area

• method

• multifidelity_polynomial_chaos

• expansion_order_sequence

Sequence of expansion orders used in a multi-stage expansion
Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One) Group 1</td>
<td></td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>expansion_samples_sequence</td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., expansion_samples.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>
7.2 method

Description

Each level entry of the expansion_order_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_order

See Also

These keywords may also be of interest:

- expansion_order

7.2.52.10.1 dimension_preference

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none

Argument(s): REALLIST

Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.52.10.2 basis_type

- Keywords Area
- method
- multifidelity_polynomial_chaos
Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

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<thead>
<tr>
<th>Required/Optional</th>
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<td>Required (Choose One)</td>
<td>PCE Basis Type (Group 1)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.

7.2.52.10.2.1 tensor_product

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- basis_type
- tensor_product

Use a tensor-product index set to construct a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none
Description

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\max (i_1, \ldots, i_d) \leq p$$

7.2.52.10.2.2 total_order

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- basis_type
- total_order

Use a total-order index set to construct a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.52.10.2.3 adapted

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

#### 7.2.52.10.2.4 advancements

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **basis_type**
- **adapted**
- **advancements**

The maximum number of steps used to expand a basis step.

### Specification

**Alias:** none

**Argument(s):** INTEGER

### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
### 7.2.52.10.2.5 soft_convergence_limit

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

### 7.2.52.10.3 collocation_ratio

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Description of Group**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional (Choose One)</td>
<td>Regression Algorithm (Group 1)</td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>-----------------------</td>
<td>--------------------------------</td>
<td>-----------------------------</td>
<td>---------------------------------------------------------------------</td>
</tr>
<tr>
<td>least_squares</td>
<td></td>
<td></td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>least_angle_regression</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
</tbody>
</table>
### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate $N$ from $n$ and $p$, the collocation_ratio allows for specification of a constant factor applied to $N$ (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with $N$ can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $cN^o \frac{n+p}{n+1}$).
7.2.52.10.3.1 collocation_points_sequence

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

**Specification**

**Alias:** pilot_samples

**Argument(s):** INTEGERLIST

**Description**

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

**See Also**

These keywords may also be of interest:

- collocation_points

7.2.52.10.3.2 least_squares

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**
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<tr>
<th>Required/Optional</th>
<th>Description of Group LSQ Regression Approach (Group 1)</th>
<th>Dakota Keyword</th>
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<tr>
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<td></td>
<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>equality_- constrained</td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.52.10.3.3 svd

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **least_squares**
- **svd**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

7.2.52.10.3.4 equality_constrained

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.52.10.3.5 orthogonal_matching_pursuit

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **orthogonal_matching_pursuit**

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.
7.2.52.10.3.6 noise_tolerance

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.52.10.3.7 basis_pursuit

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

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<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.52.10.3.9 noise_tolerance

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)
7.2 method

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.52.10.3.10 least_angle_regression

• Keywords Area
  • method
  • multifidelity_polynomial_chaos
  • expansion_order_sequence
  • collocation_ratio
  • least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.52.10.3.11 noise_tolerance

• Keywords Area
  • method
  • multifidelity_polynomial_chaos
  • expansion_order_sequence
  • collocation_ratio
  • least_angle_regression
  • noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### 7.2.52.10.3.12 least_absolute_shrinkage

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **least_absolute_shrinkage**

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td>The l2 penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.

### 7.2.52.10.3.13 noise_tolerance

- **Keywords Area**
- **method**
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

---

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

**Keyword**: noise_only

- **Keywords Area**: none
- **Argument(s)**: none
- **Description**: Restrict the cross validation process to estimating only the best noise tolerance.
Specification

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.52.10.3.17  max_cv_order_candidates

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **cross_validation**
- **max_cv_order_candidates**

Limit the number of cross-validation candidates for basis order

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

**Description**

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

**See Also**

These keywords may also be of interest:

- **increment_max_order**
- **increment_max_rank_order**
- **polynomial_chaos**
Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

**Description**

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order.

**7.2.52.10.3.19 response_scaling**

Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use original data
Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- multifidelity_polynomial_chaos
- multifidelity_function_train

7.2.52.10.3.20 use_derivatives

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it’s use with Surfpack Gaussian process is not recommended.

7.2.52.10.3.21 tensor_grid

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

**reuse_points**

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The **reuse_points** option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify **reuse_points** so that any points that have been previously generated (for example, from the import_points file) can be reused.
7.2 method

- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none

Argument(s): INTEGER

Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.52.10.4 expansion_samples_sequence

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>
See Also

These keywords may also be of interest:

- expansion_samples

7.2.52.10.4.1 reuse_points

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- expansion_samples_sequence
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.52.10.5 import_build_points_file

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

#### Default Behavior

By default, methods do not import points from a file.

#### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

#### Examples

```dakota
method
polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

**7.2.52.10.5.1 custom_annotated**

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = ‘dakota_summary.dat’
custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
</tbody>
</table>

7.2.52.10.5.2 header

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.52.10.5.3 eval_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Keywords Area

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.52.10.5.4 interface_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.52.10.5.5 annotated

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- annotated

Selects annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

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7.2.52.10.5.6 freeform

- Keywords Area
- method
- multifidelity_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
freeform

Resulting tabular file:

```
0.9     1.1     0.0002    0.26     0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
```

**7.2.52.10.5.7 active_only**

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **expansion_order_sequence**
- **import_build_points_file**
- **active_only**

Import only active variables from tabular data file

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

**7.2.52.11 orthogonal_least_interpolation**

- **Keywords Area**
- **method**
• multifidelity_polynomial_chaos
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>collocation_points-sequence</td>
<td></td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build-points-file</td>
<td></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods expansion_order cannot be set. OLI will produce the lowest degree polynomial that interpolates the data

7.2.52.11.1 collocation_points_sequence

• Keywords Area
• method
• multifidelity_polynomial_chaos
• orthogonal_least_interpolation
• collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
7.2 method

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.52.11.2 tensor_grid

- Keywords Area
- method
- multifidelity_polynomial_choa
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): INTEGERLIST
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.52.11.3 reuse_points

- Keywords Area
- method
- multifidelity_polynomial_choa
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The **reuse_points** option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

### 7.2.52.11.4 import_build_points_file

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>
Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```plaintext
method polynomial_chaos
   expansion_order = 4
   import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.52.11.4.1 custom_annotated

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias**: none

**Argument(s)**: none

**Default**: annotated format

**Child Keywords**: none
### Keywords Area

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface-_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1     0.9  1.1   0.0002 0.26 0.76
2  0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
7.2.52.11.4.2 header

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.52.11.4.3 eval_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated
7.2.52.11.4.4  interface_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom.annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom.annotated

7.2.52.11.4.5  annotated

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```bash
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.52.11.4.6 freeform

- Keywords Area
- method
- multifidelity_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.52.11.4.7 active_only
```

Import only active variables from tabular data file
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.52.12 askey

- Keywords Area
- method
- multifidelity_polynomial_chaos
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- polynomial_chaos
- askey

**normalized**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
7.2 method

Specification

Alias: none
Argument(s): none
Default: PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.52.15 export_expansion_file

- Keywords Area
- method
- multifidelity_polynomial_chaos
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

Alias: none
Argument(s): STRING

Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.52.16 samples_on_emulator

- Keywords Area
- method
- multifidelity_polynomial_chaos
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples
Argument(s): INTEGER
Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method
    polynomial_chaos
      quadrature_order = 2
      samples_on_emulator = 10000
```

7.2.52.17 sample_type

- Keywords Area
- method
- multifidelity_polynomial_chaos
- sample_type

Selection of sampling strategy

Specification

Alias: none

Argument(s): none

Default: random

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

Default Behavior
If the sample_type keyword is present, it must be accompanied by lhs or random. In most contexts, lhs is the default (exception: multilevel_sampling uses Monte Carlo by default).

Examples

method
sampling
  sample_type lhs
  samples = 20
  seed = 83921

7.2.52.17.1  lhs

  • Keywords Area
  • method
  • multifidelity_polynomial_ chaos
  • sample_type
  • lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

Alias: none
Argument(s): none

Description

The lhs keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the lhs keyword must appear in conjunction with the sample_type keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

method
  sampling
    sample_type lhs
    samples = 20

7.2.52.17.2  random

  • Keywords Area
  • method
Uses purely random Monte Carlo sampling to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```dakota
method
  sampling
    sample_type random
    samples = 200
```

**rng**

Selection of a random number generator

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**
### Description

The `rng` keyword is used to indicate a choice of random number generator.

#### Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

#### Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

#### Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.52.18.1 `mt19937`

- `Keywords Area`
- `method`
- `multifidelity_polynomial_chaos`
- `rng`
- `mt19937`

Generates random numbers using the Mersenne twister

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).
Default Behavior

mt\textsubscript{19937} is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt\textsubscript{19937}) is recommended.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
\end{verbatim}

7.2.52.18.2  \texttt{rnum2}

- Keywords Area
- method
- multifidelity\_polynomial\_chaos
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

\textbf{Alias:} none

\textbf{Argument(s):} none

Description

The \texttt{rnum2} keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

\texttt{rnum2} is not used by default. To change this behavior, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt\textsubscript{19937}) is recommended over \texttt{rnum2}.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
\end{verbatim}
7.2.52.19 probability_refinement

• Keywords Area
  • method
  • multifidelity_polynomial_chaos
  • probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

  • reliability_methods

Specification

Alias: sample_refinement

Argument(s): none

Default: no refinement

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Importance Sampling Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design</td>
<td></td>
</tr>
</tbody>
</table>

Description

The probability_refinement allows refinement of probability and generalized reliability results using importance sampling. If one specifies probability_refinement, there are some additional options. One can specify which type of importance sampling to use (import, adapt_import, or mm_adapt_import). Additionally, one can specify the number of refinement samples to use with refinement_samples and the seed to use with seed.

The probability_refinement density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.
7.2.52.19.1 import

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_refinement
- import

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

7.2.52.19.2 adapt_import

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_refinement
- adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.
7.2.52.19.3 mm_adapt_import

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_refinement
- mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

**mm_adapt_import** starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for "representative" points. Once these are located, the multimodal sampling density is set and then **mm_adapt_import** proceeds similarly to **adapt_import** (sample until convergence).

7.2.52.19.4 refinement_samples

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_refinement
- refinement_samples

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial samples in a sampling design.
7.2.52.20 final_moments

- Keywords Area
- method
- multifidelity_polynomial_chaos
- final_moments

Output moments of the specified type and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** standard

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>none</td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

**Description**

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

method, sampling
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments central

Overriding the default to none as follows:

method,
  sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none

allows associated nested model mappings to be simplified from:

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.

7.2.52.20.1 none

- Keywords Area
- method
- multifidelity_polynomial_chaos
- final_moments
- none

Omit moments from the set of final statistics.

Specification

Alias: none

Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
  sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
7.2.52.20.2 standard

- Keywords Area
- method
- multifidelity_polynomial_chaos
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none

Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments standard
```

7.2.52.20.3 central

- Keywords Area
- method
- multifidelity_polynomial_chaos
- final_moments
- central

Output central moments and include them within the set of final statistics.

Specification

Alias: none

Argument(s): none
7.2 method

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method, sampling
   samples = 50 seed = 1234
   response_levels = 3.6e+11 1.2e+05 3.5e+05
   final_moments central
```

7.2.52.21 response_levels

- Keywords Area
- method
- multifidelity_polynomial_ chaos
- response_levels

Values at which to estimate desired statistics for each response

Specification

**Alias**: none

**Argument(s)**: REALLIST

**Default**: No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior
If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```
method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
       6.0e+04 6.5e+04 7.0e+04
       3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
```

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interface,
    system async evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses,
    response_functions = 3
    no_gradients
    no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.52.21.1 num_response_levels

- Keywords Area
- method
- multifidelity_polynomial_ chaos
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

**Description**

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```
method sampling
  samples = 100
  seed = 34785
  num_response_levels = 1 1 1
  response_levels = 0.5 0.5 0.5
```

7.2.52.21.2 compute

- Keywords Area
- method
- multifidelity_polynomial_ chaos
- response_levels
- compute

Selection of statistics to compute at each response level
### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** probabilities

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td>relabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td>system</td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
</tbody>
</table>

### Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

### Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

### Expected Output

The type of statistics specified by `compute` will be reported for each response level.

### Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

### Examples

```
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary_distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```
The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

**Examples**

```verbatim
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
  6.0e+04 6.5e+04 7.0e+04
  3.5e+05 4.0e+05 4.5e+05
  compute probabilities

7.2.52.21.2.2 reliabilities
```

- **Keywords Area**
- **method**
- `multifidelity_polynomial_chaos`
- `response_levels`

Computes probabilities associated with response levels

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.
7.2 method

- compute
- reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The reliabilities keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the reliabilities are not computed by default. To change this behavior, the reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```

7.2.52.21.2.3 gen_reliabilities

- Keywords Area
- method
- multifidelity_polynomial_chaos
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none
Argument(s): none
Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities

7.2.52.21.2.4 system
  • Keywords Area
  • method
  • multifidelity_polynomial_chaos
  • response_levels
  • compute
  • system

Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>System Reliability Mode (Group 1)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.52.21.2.5 series

- Keywords Area
- method
- multifidelity_polynomial_chaos
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.52.21.2.6 parallel

- Keywords Area
- method
- multifidelity_polynomial_chaos
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system
Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.52.22 probability_levels

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Ind</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td>6.0000000000e-01</td>
<td></td>
</tr>
</tbody>
</table>
Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.52.22.1 num_probability_levels

- Keywords Area
- method
- multifidelity_polynomial_chaos
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.52.23 reliability_levels

- Keywords Area
- method
7.2 method

- `multifidelity_polynomial_chaos`
- `reliability_levels`

Specify reliability levels at which the response values will be estimated.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.52.23.1 num_reliability_levels

- `Keywords Area`  
- method  
- `multifidelity_polynomial_chaos`  
- `reliability_levels`  
- num_reliability_levels

Specify which reliability_levels correspond to which response.

**Specification**

**Alias:** none
**Argument(s):** INTEGERLIST

**Default:** reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.52.24 gen_reliability_levels

- Keywords Area
- method
- multifidelity_polynomial_chaos
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_gen_-</td>
<td>num_gen_-</td>
<td>gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
7.2.52.24.1  num_gen_reliability_levels

- Keywords Area
- method
- multifidelity_polynomial_chaos
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.52.25  distribution

- Keywords Area
- method
- multifidelity_polynomial_chaos
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
<td></td>
</tr>
</tbody>
</table>
Description

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.52.25.1 cumulative

- Keywords Area
- method
- multifidelity_polynomial_chaos
- distribution
- cumulative

Computes statistics according to cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**
Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

**7.2.52.25.2 complementary**

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **distribution**
- **complementary**

Computes statistics according to complementary cumulative functions.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

**7.2.52.26 variance_based_decomp**

- **Keywords Area**
Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics. Suppressed output of sensitivity indices with values lower than this tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>drop_tolerance</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

**Default Behavior**

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol’ indices.

**Expected Outputs**

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

**Usage Tips**

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.
Examples

```dakota
method, polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of [76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in [90].

7.2.52.26.1 interaction_order

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **variance_based_decomp**
- **interaction_order**

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

**Specification**

- **Alias:** none
- **Argument(s):** INTEGER
- **Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

**Description**

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.52.26.2 drop_tolerance

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **variance_based_decomp**
- **drop_tolerance**

Suppresses output of sensitivity indices with values lower than this tolerance.
Specification

Alias: none

Argument(s): REAL

Default: All VBD indices displayed

Description

The `drop_tolerance` keyword allows the user to specify a value below which sensitivity indices generated by `variance_based_decomp` are not displayed.

Default Behavior

By default, all sensitivity indices generated by `variance_based_decomp` are displayed.

Usage Tips

For `polynomial_chaos`, which outputs main, interaction, and total effects by default, the `univariate_effects` may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be `diagonal_covariance` or `full_covariance`, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

```
method,
  sampling
    sample_type lhs
    samples = 100
    variance_based_decomp
    drop_tolerance = 0.001
```

7.2.52.27 diagonal_covariance

- Keywords Area
- method
  - multifidelity_polynomial_chaos
  - diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none

Argument(s): none

Default: `diagonal_covariance` for response vector > 10; else `full_covariance`

Description

With a large number of responses, the covariance matrix can be very large. `diagonal_covariance` is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
7.2.52.28  full_covariance

- Keywords Area
- method
- multifidelity_polynomial_chaos
- full_covariance

Display the full covariance matrix

Specification

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.52.29  import_approx_points_file

- Keywords Area
- method
- multifidelity_polynomial_chaos
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
Keywords Area

<table>
<thead>
<tr>
<th>freeform</th>
<th>Selects freeform file format</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>active_only</td>
</tr>
<tr>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

**Default Behavior**  No import of points at which to evaluate the surrogate.

**Expected Output**  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

**Examples**

```plaintext
method
  polynomial_chaos
  expansion_order = 4
  import_approx_points_file = 'import.mcmc_annot.dat'
  annotated
```

7.2.52.29.1  custom_annotated

- **Keywords Area**
- **method**
- **multifidelity_polynomial_chaos**
- **import_approx_points_file**
- **custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
```

- Keywords Area
- method

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: no header

**Description**

See description of parent `custom_annotated`

7.2.52.29.1.2 eval_id

- **Keywords Area**
- **method**
- `multifidelity_polynomial_chaos`
- `import_approx_points_file`
- `custom_annotated`
- `eval_id`

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: no eval_id column

**Description**

See description of parent `custom_annotated`

7.2.52.29.1.3 interface_id

- **Keywords Area**
- **method**
- `multifidelity_polynomial_chaos`
- `import_approx_points_file`
- `custom_annotated`
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.52.9.2 annotated

• Keywords Area
• method
• multifidelity_polynomial_chaos
• import_approx_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

```plaintext
%eval_id interface x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  NO_ID   0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.52.29.3  freeform

• Keywords Area

• method

• multifidelity_polynomial_choas

• import_approx_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format
Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
 0.9  1.1  0.0002  0.26  0.76
 0.90009 1.1  0.0001996404857  0.2601620081  0.759955
 0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.52.29.4 active_only

- Keywords Area
- method
- multifidelity_polynomial_chaos
- import_approx_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword \texttt{active\_only} indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.52.30 \texttt{export\_approx\_points\_file}

- Keywords Area
- \texttt{method}
- \texttt{multifidelity\_polynomial\_chaos}
- \texttt{export\_approx\_points\_file}

Output file for surrogate model value evaluations

Specification

Alias: \texttt{export\_points\_file}
Argument(s): STRING
Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>annotated</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td></td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate's predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:
7.2 method

- annotated (default)
- custom.annotated
- freeform

7.2.52.30.1 custom.annotated

- Keywords Area
- method
- multifidelity.polynomial.chaos
- export.approx.points.file
- custom.annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.
Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```bash
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.52.30.1.1 header

- Keywords Area
- method
- multifidelity_polynomial_choas
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated
7.2 method

7.2.52.30.1.2 eval_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent custom_annotated

7.2.52.30.1.3 interface_id

- Keywords Area
- method
- multifidelity_polynomial_chaos
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent custom_annotated
Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated` header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.52.30.3  freeform

- Keywords Area
- method
- multifidelity_polynomial_chaos
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- fileFormats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.52.31 seed_sequence

• Keywords Area

• method

• multifidelity_polynomial_chaos

• seed_sequence

Sequence of seed values for multi-stage random sampling

Specification

Alias: none

Argument(s): INTEGERLIST

Default: system-generated (non-repeatable)

Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips
For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
   model_pointer = 'HIERARCH'
   multilevel_sampling
   pilot_samples = 100 50 40 30 20
   seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

See Also

These keywords may also be of interest:

- seed
- fixed_seed

7.2.52.32 fixed_seed

- Keywords Area
- method
- multifidelity_polynomial_chaos
- fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run

**Description**

The fixed_seed flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.
Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

7.2.52.33 model_pointer

- Keywords Area
- method
- multifidelity_polynomial_chaos
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.
Examples

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    sample_type lhs
distribution cumulative
```

```
model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic
```

```
method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2
```

```
model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'
```

```
variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'
```

```
interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'
```

```
responses
  response_functions = 3
  no_gradients
  no_hessians
```

7.2.53  multilevel_polynomial_chaos

- Keywords Area
- method
- multilevel_polynomial_chaos

Multilevel uncertainty quantification using polynomial chaos expansions

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td></td>
<td>Sample allocation approach for multilevel expansions</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td>metric_scale</td>
<td></td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td>discrepancy_-emulation</td>
<td></td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>expansion_order_-sequence</td>
<td></td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td>orthogonal_least-interpolation</td>
<td></td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>askey</td>
<td></td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td>wiener</td>
<td></td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
<tr>
<td>Optional</td>
<td>normalized</td>
<td>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>------------</td>
<td>----------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_expansion_file</td>
<td>Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate).</td>
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</tr>
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<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy.</td>
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</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value.</td>
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</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
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<tr>
<td>--------------------------------------</td>
<td>--------------------</td>
<td>--------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_-</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>levels</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_-</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>decomp</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>Covariance Type</td>
<td>diagonal_-covariance</td>
<td></td>
</tr>
<tr>
<td>Choose One</td>
<td>(Group 3)</td>
<td>Display only the diagonal terms of the covariance matrix</td>
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</tr>
<tr>
<td>Optional</td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
<td></td>
</tr>
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<td>Optional</td>
<td>import_approx_-</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate</td>
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<tr>
<td>Optional</td>
<td>points_file</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_-</td>
<td>Output file for surrogate model value evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>points_file</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td>Sequence of seed values for multi-stage random sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
<td></td>
</tr>
</tbody>
</table>
7.2 method

| Optional | model_pointer | Identifier for model block to be used by a method |

Description

As described in polynomial_chaos, the polynomial chaos expansion (PCE) is a general framework for the approximate representation of random response functions in terms of series expansions in standardized random variables:

\[ R = \sum_{i=0}^{P} \alpha_i \Psi_i(\xi) \]

where \( \alpha_i \) is a deterministic coefficient, \( \Psi_i \) is a multidimensional orthogonal polynomial and \( \xi \) is a vector of standardized random variables.

In the multilevel and multifidelity cases, we decompose this expansion into several constituent expansions, one per model form or solution control level. In a bi-fidelity case with low-fidelity (LF) and high-fidelity (HF) models, we have:

\[ R = \sum_{i=0}^{P_{LF}} \alpha_{LF,i} \Psi_i(\xi) + \sum_{i=0}^{P_{HF}} \delta_i \Psi_i(\xi) \]

where \( \delta_i \) is a coefficient for the discrepancy expansion.

For the case of regression-based PCE (least squares, compressed sensing, or orthogonal least interpolation), an optimal sample allocation procedure can be applied for the resolution of each level within a multilevel sampling procedure as in multilevel_sampling. The core difference is that a Monte Carlo estimator of the statistics is replaced with a PCE-based estimator of the statistics, requiring approximation of the variance of these estimators.

Initial prototypes for multilevel PCE can be explored using dakota/share/dakota/test/dakota_uq_-diffusion_mlpce.in, and will be stabilized in future releases.

Additional Resources

Dakota provides access to multilevel PCE methods through the NonDMultilevelPolynomialChaos class. Refer to the Stochastic Expansion Methods chapter of the Theory Manual[16] for additional information on the Multilevel PCE algorithm.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments (expansion moments only)
- Probability Density
- Level Mappings

In addition, the execution group has the attribute equiv_hf_evals, which records the equivalent number of high-fidelity evaluations.

Examples

```python
method,
    multilevel_polynomial_chaos
    model_pointer = 'HIERARCH'
    pilot_samples = 10
    expansion_order_sequence = 2
    collocation_ratio = .9
    seed = 1237
```
orthogonal_matching_pursuit
convergence_tolerance = .01
output silent

model,
   id_model = 'HIERARCH'
surrogate_hierarchical
   ordered_model_fidelities = 'SIM1'
correction_additive zeroth_order

model,
   id_model = 'SIM1'
simulation
   solution_level_control = 'mesh_size'
solution_level_cost = 1. 8. 64. 512. 4096.

See Also

These keywords may also be of interest:

- adaptive_sampling
- gpais
- local_reliability
- global_reliability
- sampling
- importance_sampling
- stoch_collocation

7.2.53.1 max_iterations

- Keywords Area
- method
- multilevel_polynomial_chaos
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt , local_reliability: 25; global_(reliability , interval_est , evidence) / efficient_global: 25∗n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.53.2 allocation_control

- Keywords Area
  - \texttt{method}
  - \texttt{multilevel\_polynomial\_chaos}
  - \texttt{allocation\_control}

Sample allocation approach for multilevel expansions

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: none

\textbf{Child Keywords}:

<table>
<thead>
<tr>
<th>Child Keyword</th>
<th>Description</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>\texttt{estimator_variance}</td>
<td>Variance of mean estimator within multilevel polynomial chaos</td>
<td>\texttt{estimator_variance}</td>
<td>Variance of mean estimator within multilevel polynomial chaos</td>
</tr>
<tr>
<td>\texttt{rip_sampling}</td>
<td>Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos</td>
<td>\texttt{rip_sampling}</td>
<td>Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

Description

Multilevel expansions, including regression-based polynomial chaos expansion (PCE) and function train (FT) expansions, require a sample allocation strategy. Three options are currently available:

- allocation based on assuming a convergence rate for the estimator variance (for regression PCE)
• restricted isometry property (RIP) sampling (for regression PCE via compressed sensing)
• rank sampling (for FT)

Default Behavior
Current defaults, when allocation_control is not specified, are estimator variance for PCE and rank sampling for FT.

7.2.53.2.1 estimator_variance

Variance of mean estimator within multilevel polynomial chaos

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>estimator_rate</td>
<td>Rate of convergence of mean estimator within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[ \text{Var}(\hat{Q}) = \frac{\sigma_Q^2}{N} \]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[ \text{Var}(\hat{Q}) = \frac{\sigma_Q^2}{\gamma N^\kappa} \]

for free parameters \( \gamma \) and \(\kappa\), with default values that may be overridden as part of this specification block.

This approach is supported for regression-based PCE approaches (over-determined least squares, under-determined compressed sensing, or orthogonal least interpolation).

In practice, it can be challenging to estimate a smooth convergence rate for estimator variance in the presence of abrupt transitions in the quality of sparse recoveries. As a result, sample allocation by greedy refinement is generally preferred.
Examples

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the convergence of the estimator variance, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```plaintext
method,
    model_pointer = 'HIERARCH'
    multilevel_polynomial_chaos
        orthogonal_matching_pursuit
        expansion_order_sequence = 2
    pilot_samples = 10
    collocation_ratio = .9
    allocation_control
        estimator_variance estimator_rate = 2.5
    seed = 1237
    convergence_tolerance = .01
```

### 7.2.53.2.1.1 estimator_rate

- **Keywords** Area
- **method**
- **multilevel_polynomial_chaos**
- **allocation_control**
- **estimator_variance**
- **estimator_rate**

Rate of convergence of mean estimator within multilevel polynomial chaos

#### Specification

**Alias:** none  
**Argument(s):** REAL  
**Default:** 2

#### Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}
\]

for free parameters \(\gamma\) and \(\kappa\).

The default values are \(\gamma = 1\) and \(\kappa = 2\) (adopts a more aggressive sample profile by assuming a faster convergence rate than Monte Carlo). This advanced specification option allows to user to specify \(\kappa\), overriding the default.
7.2.53.2.2 rip_sampling

- Keywords Area
- method
- multilevel_polynomial_chaos
- allocation_control
- rip_sampling

Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Multilevel polynomial chaos with compressed sensing may allocate the number of samples per level based on the restricted isometry property (RIP), applied to recovery at level \( l \):

\[
N_l \geq s_l \log^3(s_l) L_l \log(C_l)
\]

for sparsity \( s \), cardinality \( C \), and mutual coherence \( L \). The adaptive algorithm starts from a pilot sample, shapes the profile based on observed sparsity, and iterates until convergence. In practice, RIP sampling levels are quite conservative, and a collocation ratio constraint (\( N_l \leq rC_l \), where \( r \) defaults to 2) must be enforced on the profile.

The algorithm relies on observed sparsity, it is appropriate for use with regularized solvers for compressed sensing. It employs orthogonal matching pursuit (OMP) by default and automatically activates cross-validation in order to choose the best noise parameter value for the recovery.

This capability is **experimental**. Sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the number of sparse coefficient sets recovered for each level, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```plaintext
method,
    model_pointer = 'HIERARCH'
    multilevel_polynomial_chaos
        orthogonal_matching_pursuit
        expansion_order_sequence = 2
    pilot_samples = 10
    collocation_ratio = .9
    allocation_control rip_sampling
    seed = 1237
    convergence_tolerance = .01
```

7.2.53.3 convergence_tolerance

- Keywords Area
- method
7.2 method

- multilevel_polynomial_chaos
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do. Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.53.4 metric_scale

- Keywords Area
- method
- multilevel_polynomial_chaos
- metric_scale

define scaling of statistical metrics when adapting UQ surrogates
## Specification

**Alias:** none  
**Argument(s):** none  
**Default:** relative

### Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group scale of refinement metric (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>absolute</td>
<td>relative</td>
<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>absolute</td>
</tr>
</tbody>
</table>

### Description

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The `absolute` option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The `relative` option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

The default is `relative`.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.53.4.1 `relative`

- **Keywords Area**

- **method**

- **multilevel_polynomial_chaos**

- **metric_scale**

- **relative**

relative scaling of statistical metrics when adapting UQ surrogates
Description

When adapting a UQ surrogate model, the relative option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.53.4.2 absolute

- Keywords Area
- method
- multilevel_polynomial_chaos
- metric_scale
- absolute

use of absolute statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none

Description

When adapting a UQ surrogate model, the absolute option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

Default Behavior

Expected Output

Usage Tips

Additional Discussion

7.2.53.5 discrepancy_emulation

- Keywords Area
- method
- multilevel_polynomial_chaos
- discrepancy_emulation

Formulation for emulation of model discrepancies.
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** distinct

### Child Keywords:

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<th>Required/Optional Required (Choose One)</th>
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<td>distinct</td>
<td>Distinct formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>recursive</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

### Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.

### Examples

```method
  multilevel_polynomial_chaos
  expansion_order_sequence = 2
  collocation_ratio = .9
  orthogonal_matching_pursuit
  discrepancy_emulation recursive
```

**7.2.53.5.1 distinct**

- Keywords Area
- method
- multilevel_polynomial_chaos
- discrepancy_emulation
7.2 method

- distinct

Distinct formulation for emulation of model discrepancies.

Specification

Alias: paired

Argument(s): none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.53.5.2 recursive

- Keywords Area
- method
- multilevel_polynomial_chaos
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none

Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.53.6 expansion_order_sequence

- Keywords Area
- method
• multilevel_polynomial_chaos

• expansion_order_sequence

Sequence of expansion orders used in a multi-stage expansion

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:
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<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_samples_sequence</td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., expansion_samples.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>
Description

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

See Also

These keywords may also be of interest:

- `expansion_order`

7.2.53.6.1 `dimension_preference`

- `Keywords Area`
- `method`
- `multilevel_polynomial_chaos`
- `expansion_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension).

Specification

**Alias**: none

**Argument(s)**: REALLIST

**Default**: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- `sobol`
- `decay`

7.2.53.6.2 `basis_type`

- `Keywords Area`
- `method`
- `multilevel_polynomial_chaos`
Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>PCE Basis Type (Group 1)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

**Description**

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.

7.2.53.6.2.1 tensor_product

- **Keywords Area**
  - method
  - multilevel_polynomial_chaos
  - expansion_order_sequence
  - basis_type
  - tensor_product

Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none
Description

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\max (i_1, \ldots, i_d) \leq p$$

7.2.53.6.2.2 total_order

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- basis_type
- total_order

Use a total-order index set to construct a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none

Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.53.6.2.3 adapted

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

#### 7.2.53.6.2.4 advancements

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **expansion_order_sequence**
- **basis_type**
- **adapted**
- **advancements**

The maximum number of steps used to expand a basis step.

### Specification

**Alias:** none

**Argument(s):** INTEGER

### Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.53.6.2.5 soft_convergence_limit

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.

Specification

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.53.6.3 collocation_ratio

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

**Alias:** none

**Argument(s):** REAL

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
<table>
<thead>
<tr>
<th>Optional</th>
<th>Regression Algorithm (Group 1)</th>
<th>collocation_points-sequence</th>
<th>Sequence of collocation point counts used in a multi-stage expansion</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td></td>
<td>orthogonal_matching_pursuit</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
<td></td>
<td>basis_pursuit</td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
</tr>
<tr>
<td></td>
<td>basis_pursuit_denoising</td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td></td>
<td>least_angle_regression</td>
<td>least_angle_regression</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td></td>
<td>least_absolute_shrinkage</td>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
<tr>
<td>Optional</td>
<td>cross_validation</td>
<td>Use cross validation to choose the ‘best’ polynomial order of a polynomial chaos expansion. Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>ratio_order</td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients.</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $cN^o_{n+1}$).
7.2.53.6.3.1 collocation_points_sequence

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

**Specification**

**Alias:** pilot_samples

**Argument(s):** INTEGERLIST

**Description**

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

**See Also**

These keywords may also be of interest:

- collocation_points

7.2.53.6.3.2 least_squares

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**
### Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

**7.2.53.6.3.3 svd**

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **least_squares**
- **svd**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

### Specification

**Alias:** none

**Argument(s):** none

### Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

**7.2.53.6.3.4 equality_constrained**

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
7.2 method

- expansion_order_sequence
- collocation_ratio
- least_squares
- equality_constrained

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.53.6.3.5 orthogonal_matching_pursuit

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.
7.2.53.6.3.6 noise_tolerance

• Keywords Area
• method
• multilevel_polynomial_chaos
• expansion_order_sequence
• collocation_ratio
• orthogonal_matching_pursuit
• noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none
Argument(s): REALLIST
Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.53.6.3.7 basis_pursuit

• Keywords Area
• method
• multilevel_polynomial_chaos
• expansion_order_sequence
• collocation_ratio
• basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

Specification

Alias: bp
Argument(s): none

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
7.2.53.6.3.8 basis_pursuit_denoising

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
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<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.53.6.3.9 noise_tolerance

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)
Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.53.6.3.10 least_angle_regression

- Keywords Area
  - method
  - multilevel_polynomial_chaos
  - expansion_order_sequence
  - collocation_ratio
  - least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.53.6.3.11 noise_tolerance

- Keywords Area
  - method
  - multilevel_polynomial_chaos
  - expansion_order_sequence
  - collocation_ratio
  - least_angle_regression
  - noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.53.6.3.12 least_absolute_shrinkage

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso

Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td></td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.53.6.3.13 noise_tolerance

- Keywords Area
- method
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** \(1e-3\) for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

---

The \(l_2\) penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The \(l_2\) penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

<table>
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<tr>
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<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_only</td>
<td>max_cv_order_ candidates</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance. Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

**7.2.53.6.3.16 noise_only**

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **expansion_order_sequence**
- **collocation_ratio**
- **cross_validation**

Restrict the cross validation process to estimating only the best noise tolerance.
Specification

Alias: none
Argument(s): none
Default: false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.53.6.3.17 max_cv_order_candidates

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
7.2.53.6.3.18 ratio_order

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

**Description**

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $c N^{\omega}$ where $c$ is the collocation_ratio and $\omega$ is the ratio_order.

7.2.53.6.3.19 response_scaling

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use original data
Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.53.6.3.20 `use_derivatives`

- `Keywords Area`
- `method`
- `multilevel_polynomial_chaos`
- `expansion_order_sequence`
- `collocation_ratio`
- `use_derivatives`

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.53.6.3.21 `tensor_grid`

- `Keywords Area`
- `method`
- `multilevel_polynomial_chaos`
- `expansion_order_sequence`
- `collocation_ratio`
7.2 method

- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.53.6.3.22 reuse_points

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.53.6.3.23 max_solver_iterations

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
Keywords Area

• collocation_ratio
• max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.53.6.4 expansion_samples_sequence

• Keywords Area
• method
• multilevel_polynomial_chaos
• expansion_order_sequence
• expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

Specification

Alias: none
Argument(s): INTEGERLIST
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>
See Also

These keywords may also be of interest:

- expansion_samples

7.2.53.6.4.1 reuse_points

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- expansion_samples_sequence
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.53.6.5 import_build_points_file

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

Child Keywords:
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```plaintext
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.53.6.5.1 custom_annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom.annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.53.6.5.2 header

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.53.6.5.3 eval_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
### 7.2 method

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

#### Description

See description of parent `custom_annotated`

**7.2.53.6.5.4 interface_id**

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **expansion_order_sequence**
- **import_build_points_file**
- **custom_annotated**
- **interface_id**

 Enable interface ID column in custom-annotated tabular file

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

#### Description

See description of parent `custom_annotated`

**7.2.53.6.5.5 annotated**

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **expansion_order_sequence**
- **import_build_points_file**
- **annotated**

Selects annotated tabular file format
Topics
This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description
An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior
By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
      annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
7.2.53.6.5.6  freeform

- Keywords Area
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9   1.1   0.0002  0.26   0.76
0.90009 1.1   0.0001996404857  0.2601620081  0.759955
0.89991 1.1   0.0002003604863  0.2598380081  0.760045
...
```

7.2.53.6.5.7 active_only

- Keywords Area
- method
- multilevel_polynomial_ chaos
- expansion_order_sequence
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none  
**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.53.7 orthogonal_least_interpolation

- Keywords Area
- method
• multilevel_polynomial_chaos
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>collocation_points_sequence</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build_points_file</td>
<td></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods expansion_order cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

7.2.53.7.1 collocation_points_sequence

• Keywords Area
• method
• multilevel_polynomial_chaos
• orthogonal_least_interpolation
• collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
Specification

Alias: none
Argument(s): INTEGERLIST

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.53.7.2 tensor_grid

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): INTEGERLIST
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.53.7.3 reuse_points

- Keywords Area
- method
- multilevel_polynomial_chaos
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

### 7.2.53.7.4 import_build_points_file

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Optional | active_only | Import only active variables from tabular data file |
Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```method
dakota
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.53.7.4.1 custom_annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1   x2   obj_fn  nln_ineq_con_1 nln_ineq_con_2
1         0.9  1.1   0.0002  0.26         0.76
2        0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3        0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
7.2.53.7.4.2 header

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

7.2.53.7.4.3 eval_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent custom_annotated
7.2.53.7.4.4 interface_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.53.7.4.5 annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```dakota
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

%eval_id interface x1  x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID    0.9 1.1  0.0002  0.26  0.76
2   NO_ID   0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID   0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.53.7.4.6 freeform

- Keywords Area
- method
- multilevel_polynomial_chaos
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

environment
tabular_data
  tabular_data_file = ‘dakota_summary.dat’
  freeform

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.9009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.00020003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.53.7.4.7 active_only

• Keywords Area

• method

• multilevel_polynomial_chaos

• orthogonal_least_interpolation

• import_build_points_file

• active_only

Import only active variables from tabular data file
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.53.8 askey

- Keywords Area
- method
- multilevel_polynomial_chaos
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).
7.2 method 1427

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

7.2.53.9 wiener

- Keywords Area
- method
- multilevel_polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.53.10 normalized

- Keywords Area
- method
- multilevel_polynomial_chaos
- normalized

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials
Keyword: export_expansion_file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

**Description**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

Keyword: samples_on_emulator

**Specification**

**Alias:** samples

**Argument(s):** INTEGER

**Description**

Number of samples at which to evaluate an emulator (surrogate).
Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
    polynomial_chaos
        quadrature_order = 2
        samples_on_emulator = 10000
```

7.2.5.3.13 sample_type

- Keywords Area
- method
- multilevel_polynomial_chaos
- sample_type

Selection of sampling strategy

Specification

**Alias:** none

**Argument(s):** none

**Default:** random

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

Default Behavior
If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

7.2.53.13.1 `lhs`
- Keywords Area
- method
- multilevel_polynomial_chaos
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

- **Alias**: none
- **Argument(s)**: none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.53.13.2 `random`
- Keywords Area
- method
7.2 method

- multilevel_polynomial_chaos
- sample_type
- random

Uses purely random Monte Carlo sampling to sample variables

Specification

Alias: none
Argument(s): none

Description

The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

method sampling
    sample_type random
    samples = 200

7.2.53.14rng

- Keywords Area
- method
- multilevel_polynomial_chaos
- rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none
Default: Mersenne twister (mt19937)
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.53.14.1 mt19937

- Keywords Area
- method
- multilevel_polynomial_chaos
- rng
- mt19937

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).
Default Behavior

\texttt{mt19937} is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended.

Examples

\begin{verbatim}
method
sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng mt19937
\end{verbatim}

7.2.53.14.2 \texttt{rnum2}

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{multilevel_polynomial_chaos}
- \texttt{rng}
- \texttt{rnum2}

Generates pseudo-random numbers using the Pecos package

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: none

Description

The \texttt{rnum2} keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

\texttt{rnum2} is not used by default. To change this behavior, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended over \texttt{rnum2}.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
\end{verbatim}
### Keywords Area

- **Keyword**: probability_refinement
  - **Description**: Allows refinement of probability and generalized reliability results using importance sampling.

### Topics

This keyword is related to the topics:

- **reliability_methods**

### Specification

**Alias**: sample_refinement

**Argument(s)**: none

**Default**: no refinement

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Importance Sampling Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Required (Choose One)</td>
<td>Importance sampling option for probability refinement</td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design.</td>
<td></td>
</tr>
</tbody>
</table>

### Description

The **probability_refinement** allows refinement of probability and generalized reliability results using importance sampling. If one specifies **probability_refinement**, there are some additional options. One can specify which type of importance sampling to use (**import**, **adapt_import**, or **mm_adapt_import**). Additionally, one can specify the number of refinement samples to use with **refinement_samples** and the seed to use with **seed**.

The **probability_refinement** density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.
7.2 method

7.2.53.1 import

• Keywords Area
• method
• multilevel_polynomial_chaos
• probability_refinement
• import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

7.2.53.2 adapt_import

• Keywords Area
• method
• multilevel_polynomial_chaos
• probability_refinement
• adapt_import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.
### mm_adapt_import

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **probability_refinement**
- **mm_adapt_import**

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

```
mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).
```

### refinement_samples

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **probability_refinement**
- **refinement_samples**

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.
7.2 final_moments

- Keywords Area
  - method
  - multilevel_polynomial_choas
  - final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td>standard</td>
<td></td>
<td></td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td>central</td>
<td></td>
<td></td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

method, sampling
overrides the default to none as follows:

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central

allows associated nested model mappings to be simplified from:

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.

7.2.53.16.1 none

- Keywords Area
- method
- multilevel_polynomial_chaos
- final_moments
- none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
7.2.53.16.2  standard

- Keywords Area
- method
- multilevel_polynomial_chaos
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments standard
```

7.2.53.16.3  central

- Keywords Area
- method
- multilevel_polynomial_chaos
- final_moments
- central

Output central moments and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none
Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```dakota
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

7.2.53.17 response_levels

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **response_levels**

Values at which to estimate desired statistics for each response

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior
If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

Examples

For example, specifying a `response_level` of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```
method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
```
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.874233192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.53.17.1 num_response_levels

- Keywords Area
- method
- multilevel_polynomial_chaos
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5

7.2.53.17.2 compute
```

Selection of statistics to compute at each response level
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** probabilities

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td><strong>probabilities</strong></td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><strong>reliabilities</strong></td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>gen_reliabilities</strong></td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>system</strong></td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method sampling  
sample_type random  
samples = 100 seed = 1  
complementary distribution  
response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```
### 7.2.53.17.2.1 probabilities

- **Keywords Area**
- **method**
- **multilevel_polynomial_chaos**
- **response_levels**
- **compute**
- **probabilities**

Computes probabilities associated with response levels

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

#### Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

#### Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

#### Examples

```
method sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
        6.0e+04 6.5e+04 7.0e+04
        3.5e+05 4.0e+05 4.5e+05
    compute probabilities
```
• compute
• reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The reliabilities keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the reliabilities are not computed by default. To change this behavior, the reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.

Examples

method sampling
   sample_type random
   samples = 100 seed = 1
   complementary distribution
   response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
   compute reliabilities

7.2.53.17.2.3 gen_reliabilities

• Keywords Area
• method
• multilevel_polynomial_choas
• response_levels
• compute
• gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none
Argument(s): none
Description

The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

7.2.53.17.2.4 system

- Keywords Area
- method
- multilevel_polynomial_choas
- response_levels
- compute
- system

Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required(Choose One)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
<td></td>
</tr>
<tr>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
<td></td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.53.17.2.5 series

• Keywords Area
• method
• multilevel_polynomial_chaos
• response_levels
• compute
• system
• series

Aggregate response statistics assuming a series system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.53.17.2.6 parallel

• Keywords Area
• method
• multilevel_polynomial_chaos
• response_levels
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none
7.2 method

Description

See parent keyword system for description.

7.2.53.18 probability_levels

- Keywords Area
- method
- multilevel_polynomial_choas
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```plaintext
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
  1. .8 .5 0.
  1. .3 .2 0.
```
variables,
   normal_uncertain = 2
   means = 248.89, 593.33
   std_deviations = 12.4, 29.7
   descriptors = ‘TF1n’ ‘TF2n’
uniform_uncertain = 2
   lower_bounds = 199.3, 474.63
   upper_bounds = 298.5, 712.
   descriptors = ‘TF1u’ ‘TF2u’
weibull_uncertain = 2
   alphas = 12., 30.
   betas = 250., 590.
   descriptors = ‘TF1w’ ‘TF2w’
histogram_bin_uncertain = 2
   num_pairs = 3 4
   abscissas = 5 8 10 .1 .2 .3 .4
   counts = 17 21 0 12 24 12 0
   descriptors = ‘TF1h’ ‘TF2h’
histogram_point_uncertain
   real = 1
   num_pairs = 2
   abscissas = 3 4
   counts = 1 1
   descriptors = ‘TF3h’

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
   Bin Lower   Bin Upper   Density Value
       ---------   ---------   --------------
2.7604749078e+11 3.4221494996e+11 5.1384774972e-12
3.4221494996e+11 4.0634975300e+11 5.1454122311e-12
4.0634975300e+11 5.4196114379e+11 2.4334239039e-12

PDF for response_fn_2:
   Bin Lower   Bin Upper   Density Value
       ---------   ---------   --------------
4.6431154744e+04 5.6511827775e+04 5.1384774972e-12
5.6511827775e+04 6.1603813790e+04 5.1454122311e-12
6.1603813790e+04 7.8702465755e+04 2.4334239039e-12

PDF for response_fn_3:
   Bin Lower   Bin Upper   Density Value
       ---------   ---------   --------------
2.3796373090e+05 3.6997214133e+05 5.1384774972e-12
3.6997214133e+05 3.8100966235e+05 5.1454122311e-12
3.8100966235e+05 4.4111498127e+05 2.4334239039e-12

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
   Response Level   Probability Level   Reliability Index   General Rel Index
                     --------------   -----------------   -----------------
2.7604749078e+11 1.0000000000e+00
3.4221494996e+11 8.0000000000e-01
4.0634975300e+11 6.0000000000e-01
5.4196114379e+11 4.0000000000e-01

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
   Response Level   Probability Level   Reliability Index   General Rel Index
                     --------------   -----------------   -----------------
4.6431154744e+04 1.0000000000e+00
5.6511827775e+04 8.0000000000e-01

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

### Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

#### 7.2.53.18.1\ num\_probability\_levels

- **Keywords Area**
- **method**
- **multilevel\_polynomial\_chaos**
- **probability\_levels**
- **num\_probability\_levels**

Specify which **probability\_levels** correspond to which response

#### Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: probability\_levels evenly distributed among response functions

#### Description

See parent page

#### 7.2.53.19\ reliability\_levels

- **Keywords Area**
- **method**
- **multilevel\_polynomial\_chaos**
- **reliability\_levels**

Specify reliability levels at which the response values will be estimated
Specification

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td>reliability_levels_correspond_to_response</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.53.19.1 num_reliability_levels

- Keywords Area
- method
- multilevel_polynomial_chaos
- reliability_levels
- num_reliability_levels

Specify which reliability_levels correspond to which response

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** reliability_levels evenly distributed among response functions

**Description**

See parent page
7.2.53.20 gen_reliability_levels

• Keywords Area
• method
• multilevel_polynomial_chaos
• gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
Specification

Alias: none
Argument(s): INTEGERLIST
Default: gen_reliability_levels evenly distributed among response functions

Description
See parent page

7.2.53.21 distribution

- Keywords Area
- method
- multilevel_polynomial_chaos
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none
Argument(s): none
Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs
Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.53.21.1 cumulative

- **Keywords Area**
- **method**
- **multilevel_polynomial_choas**
- **distribution**
- **cumulative**

Computes statistics according to cumulative functions

Specification

**Alias**: none

**Argument(s)**: none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.53.21.2 complementary

- **Keywords Area**
- **method**
- **multilevel_polynomial_choas**
• distribution
• complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

method
  sampling
  sample_type lhs
  samples = 10
  distribution complementary

7.2.53.22  variance_based_decomp

• Keywords Area
• method
• multilevel_polynomial_chaos
• variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

Alias: none
Argument(s): none
Default: no variance-based decomposition
Child Keywords:
### Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

### Default Behavior

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol’ indices.

### Expected Outputs

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

### Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.

### Examples

```method,
multiplic Polinomial_ chaos # or stoch_collocation
sparse_grid_level = 3
variance_based_decomp interaction_order = 2```

### Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76] : "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the
input variable i has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].

7.2.53.22.1 interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

**Description**

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume)

7.2.53.22.2 drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed
Description

The `drop_tolerance` keyword allows the user to specify a value below which sensitivity indices generated by `variance_based_decomp` are not displayed.

Default Behavior

By default, all sensitivity indices generated by `variance_based_decomp` are displayed.

Usage Tips

For `polynomial_chaos`, which outputs main, interaction, and total effects by default, the `univariate_effects` may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be `diagonal_covariance` or `full_covariance`, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

```dakota
method,
sampling
   sample_type lhs
   samples = 100
   variance_based_decomp
drop_tolerance = 0.001
```

7.2.53.23 diagonal_covariance

- Keywords Area
- method
- multilevel_polynomial_chaos
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none

Argument(s): none

Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. `diagonal_covariance` is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.53.24 full_covariance

- Keywords Area
- method
- multilevel_polynomial_chaos
• full_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.

7.2.53.25 import_approx_points_file

• Keywords Area
• method
• multilevel_polynomial_chaos
• import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

Specification

Alias: none
Argument(s): STRING
Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Optional/Choose One</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword custom_annotated</th>
<th>Dakota Keyword Description Selects custom-annotated tabular file format</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Default Behavior No import of points at which to evaluate the surrogate.
Expected Output  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) pro-
vided in the file and results tabulated and/or statistics computed at them, depending on the method context.

Examples

method
  polynomial_chaos
    expansion_order = 4
    import_approx_points_file = 'import.mcmc_annot.dat'

7.2.53.25.1  custom_annotated

• Keywords Area
• method
• multilevel_polynomial_chaos
• import_approx_points_file
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias:  none
Argument(s):  none
Default:  annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
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<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```dakota
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

Specification

Alias: none
Argument(s): none
Default: no header
7.2 method

Description

See description of parent custom_annotated

7.2.53.25.1.2 eval_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- import_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.53.25.1.3 interface_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- import_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated
7.2.53.25.2 annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- import_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
Resulting tabular file:

```
 eval_id interface  x1   x2    obj_fn  nln_ineq_con_1 nln_ineq_con_2
 1  NO_ID     0.9  1.1  0.0002   0.26      0.76
 2  NO_ID     0.90009 1.1 0.0001996404857 0.2601620081 0.759955
 3  NO_ID     0.89991 1.1 0.0002003604863 0.2598380081 0.760045
```

7.2.53.25.3 freeform

- Keywords Area
- method
- multilevel_polynomial_chaos
- import_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9     1.1     0.0002    0.26     0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.7599555
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.53.25.4 active_only

• Keywords Area

• method

• multilevel_polynomial_chaos

• import_approx_points_file

• active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
7.2.53.26 export_approx_points_file

- Keywords Area
- method
- multilevel_polynomial_chaos
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.53.26.1 custom_annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format
This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom.annotated header eval_id

Resulting tabular file:
% eval_id x1 x2 obj_fn nln.ineq_con_1 nln.ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...

7.2.53.26.1.1 header

- Keywords Area
- method
- multilevel_polynomial.chaos
- export.approx.points_file
- custom.annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom.annotated

7.2.53.26.1.2 eval_id

- Keywords Area
- method
- multilevel_polynomial.chaos
- export.approx.points_file
- custom.annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column
Description

See description of parent custom_annotated

7.2.53.26.1.3 interface_id

- Keywords Area
- method
- multilevel_polynomial_chaos
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no interface_id column

Description

See description of parent custom_annotated

7.2.53.26.2 annotated

- Keywords Area
- method
- multilevel_polynomial_chaos
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
@eval_id interface  x1   x2  obj_fn  nln_ineq_con_l  nln_ineq_con_2
1   NO_ID    0.9   1.1   0.0002 0.26 0.76
2   NO_ID    0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID    0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.53.26.3 freeform

- Keywords Area

- method

- multilevel_polynomial_chaos

- export_approx_points_file

- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.53.27 seed_sequence

- Keywords Area
- method
- multilevel_polynomial_chaos
- seed_sequence

Sequence of seed values for multi-stage random sampling
Specification

Alias: none

Argument(s): INTEGERLIST

Default: system-generated (non-repeatable)

Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 100 50 40 30 20
  seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

See Also

These keywords may also be of interest:

- seed
- fixed_seed

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Reuses the same seed value for multiple random sampling sets

Specification

Alias: none

Argument(s): none

Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed

Topics

This keyword is related to the topics:

- block_pointer
Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
```
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

### 7.2.54 stoch_collocation

- **Keywords Area**
  - method
  - stoch_collocation

Uncertainty quantification with stochastic collocation

**Specification**

**Alias:** nond_stoch_collocation

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td><strong>Optional (Choose One)</strong></td>
<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>h_refinement</td>
<td>Employ h-refinement to refine the grid</td>
</tr>
<tr>
<td></td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td></td>
<td></td>
<td>metric_scale</td>
<td>Define scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
<tr>
<td><strong>Required (Choose One)</strong></td>
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<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
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<td>Basis Polynomial Family (Group 3)</td>
<td>use_derivatives</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
</tr>
<tr>
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<td>----------------------------------</td>
<td>----------------</td>
<td>------------------------------------------------------------------</td>
</tr>
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<td>sample_type</td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>probability_refinement</td>
<td>Selection of a random number generator</td>
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<td>Optional</td>
<td>sparse_grid_level</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
</tr>
<tr>
<td>Optional</td>
<td>askey</td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
<tr>
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<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>--------------</td>
<td>------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
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<td>Specify probability levels at which to estimate the corresponding response value.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects.</td>
<td></td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 4)</td>
<td>diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix.</td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate.</td>
<td></td>
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</table>
### Description

Stochastic collocation is a general framework for approximate representation of random response functions in terms of finite-dimensional interpolation bases.

The stochastic collocation (SC) method is very similar to polynomial_chaos, with the key difference that the orthogonal polynomial basis functions are replaced with interpolation polynomial bases. The interpolation polynomials may be either local or global and either value-based or gradient-enhanced. In the local case, valued-based are piecewise linear splines and gradient-enhanced are piecewise cubic splines, and in the global case, valued-based are Lagrange interpolants and gradient-enhanced are Hermite interpolants. A value-based expansion takes the form

\[
R = \sum_{i=1}^{N_p} r_i L_i(\xi)
\]

where \(N_p\) is the total number of collocation points, \(r_i\) is a response value at the \(i^{th}\) collocation point, \(L_i\) is the \(i^{th}\) multidimensional interpolation polynomial, and \(\xi\) is a vector of standardized random variables.

Thus, in PCE, one forms coefficients for known orthogonal polynomial basis functions, whereas SC forms multidimensional interpolation functions for known coefficients.

#### Basis polynomial family (Group 2)

In addition to the askey and wiener basis types also supported by polynomial_chaos, SC supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis options provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space. The global gradient-enhanced interpolants (Hermite interpolation polynomials) are also restricted to uniform or transformed uniform random variables (due to the need to compute collocation weights by integration of the basis polynomials) and share the variable support shown in variable_support for Piecewise SE. Due to numerical instability in these high-order basis polynomials, they are deactivated by default but can be activated by developers using a compile-time switch.

#### Interpolation grid type (Group 3)

To form the multidimensional interpolants \(L_i\) of the expansion, two options are provided.

1. interpolation on a tensor-product of Gaussian quadrature points (specified with quadrature_order and,
optionally, \texttt{dimension\_preference} for anisotropic tensor grids). As for PCE, non-nested Gauss rules are employed by default, although the presence of \texttt{p\_refinement} or \texttt{h\_refinement} will result in default usage of nested rules for normal or uniform variables after any variable transformations have been applied (both defaults can be overridden using explicit \texttt{nested} or \texttt{non\_nested} specifications).

2. interpolation on a Smolyak sparse grid (specified with \texttt{sparse\_grid\_level} and, optionally, \texttt{dimension\_preference} for anisotropic sparse grids) defined from Gaussian rules. As for sparse PCE, nested rules are employed unless overridden with the \texttt{non\_nested} option, and the growth rules are restricted unless overridden by the \texttt{unrestricted} keyword.

Another distinguishing characteristic of stochastic collocation relative to \texttt{polynomial\_chaos} is the ability to reformulate the interpolation problem from a nodal interpolation approach into a hierarchical formulation in which each new level of interpolation defines a set of incremental refinements (known as hierarchical surpluses) layered on top of the interpolants from previous levels. This formulation lends itself naturally to uniform or adaptive refinement strategies, since the hierarchical surpluses can be interpreted as error estimates for the interpolant. Either global or local/piecewise interpolants in either value-based or gradient-enhanced approaches can be formulated using hierarchical interpolation. The primary restriction for the hierarchical case is that it currently requires a sparse grid approach using nested quadrature rules (Genz-Keister, Gauss-Patterson, or Newton-Cotes for standard normals and standard uniforms in a transformed space: Askey, Wiener, or Piecewise settings may be required), although this restriction can be relaxed in the future. A selection of hierarchical interpolation will provide greater precision in the increments to mean, standard deviation, covariance, and reliability-based level mappings induced by a grid change within uniform or goal-oriented adaptive refinement approaches (see following section).

It is important to note that, while \texttt{quadrature\_order} and \texttt{sparse\_grid\_level} are array inputs, only one scalar from these arrays is active at a time for a particular expansion estimation. These scalars can be augmented with \texttt{dimension\_preference} to support anisotropy across the random dimension set. The array inputs are present to support advanced use cases such as multifidelity UQ, where multiple grid resolutions can be employed.

\textbf{Automated refinement type (Group 1)}

Automated expansion refinement can be selected as either \texttt{p\_refinement} or \texttt{h\_refinement}, and either refinement specification can be either \texttt{uniform} or \texttt{dimension\_adaptive}. The \texttt{dimension\_adaptive} case can be further specified as either \texttt{sobol} or \texttt{generalized} (decay not supported). Each of these automated refinement approaches makes use of the \texttt{max\_iterations} and \texttt{convergence\_tolerance} iteration controls. The \texttt{h\_refinement} specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the \texttt{piecewise} specification option (it is not necessary to redundantly specify \texttt{piecewise} in the case of \texttt{h\_refinement}). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current \texttt{uniform} and \texttt{dimension\_adaptive} options.

\textbf{Covariance type (Group 5)}

These two keywords are used to specify how this method computes, stores, and outputs the covariance of the responses. In particular, the diagonal covariance option is provided for reducing post-processing overhead and output volume in high dimensional applications.

\textbf{Active Variables}

The default behavior is to form expansions over aleatory uncertain continuous variables. To form expansions over a broader set of variables, one needs to specify \texttt{active} followed by \texttt{state\_epistemic\_design\_all} in the variables specification block.

For continuous design, continuous state, and continuous epistemic uncertain variables included in the expansion, interpolation points for these dimensions are based on Gauss-Legendre rules if non-nested, Gauss-Patterson rules if nested, and Newton-Cotes points in the case of piecewise bases. Again, when probability integrals are evaluated, only the aleatory random variable domain is integrated, leaving behind a polynomial relationship between the statistics and the remaining design/state/epistemic variables.

\textbf{Optional Keywords regarding method outputs}

Each of these sampling specifications refer to sampling on the SC approximation for the purposes of generating approximate statistics.

- \texttt{sample\_type}
- \texttt{samples}
Since SC approximations are formed on structured grids, there should be no ambiguity with simulation sampling for generating the SC expansion.

When using the probability_refinement control, the number of refinement samples is not under the user's control (these evaluations are approximation-based, so management of this expense is less critical). This option allows for refinement of probability and generalized reliability results using importance sampling.

Multi-fidelity UQ

When using multifidelity UQ, the high fidelity expansion generated from combining the low fidelity and discrepancy expansions retains the polynomial form of the low fidelity expansion (only the coefficients are updated). Refer to polynomial_chaos for information on the multifidelity interpretation of array inputs for quadrature_order and sparse_grid_level.

Usage Tips

If \( n \) is small, then tensor-product Gaussian quadrature is again the preferred choice. For larger \( n \), tensor-product quadrature quickly becomes too expensive and the sparse grid approach is preferred. For self-consistency in growth rates, nested rules employ restricted exponential growth (with the exception of the dimension_adaptive p-refinement generalized case) for consistency with the linear growth used for non-nested Gauss rules (integrand precision \( i = 4l + 1 \) for sparse grid level \( l \) and \( i = 2m - 1 \) for tensor grid order \( m \)).

Additional Resources


Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments
- Probability Density
- Level Mappings
- Variance-Based Decomposition (Sobol' Indices)

Examples

```dakota
method,
  stoch_collocation
    sparse_grid_level = 2
    samples = 10000 seed = 12347 rng rnum2
    response_levels = .1 1. 50. 100. 500. 1000.
    variance_based_decomp
```
Theory

As mentioned above, a value-based expansion takes the form

\[ R = \sum_{i=1}^{N_p} r_i L_i(\xi) \]

The \( i^{th} \) interpolation polynomial assumes the value of 1 at the \( i^{th} \) collocation point and 0 at all other collocation points, involving either a global Lagrange polynomial basis or local piecewise splines. It is easy to see that the approximation reproduces the response values at the collocation points and interpolates between these values at other points. A gradient-enhanced expansion (selected via the `use_derivatives` keyword) involves both type 1 and type 2 basis functions as follows:

\[ R = \sum_{i=1}^{N_p} [r_i H_i^{(1)}(\xi) + \frac{d}{d\xi} H_i^{(2)}(\xi)] \]

where the \( i^{th} \) type 1 interpolant produces 1 for the value at the \( i^{th} \) collocation point, 0 for values at all other collocation points, and 0 for derivatives (when differentiated) at all collocation points, and the \( i,j^{th} \) type 2 interpolant produces 0 for values at all collocation points, 1 for the \( j^{th} \) derivative component at the \( i^{th} \) collocation point, and 0 for the \( j^{th} \) derivative component at all other collocation points. Again, this expansion reproduces the response values at each of the collocation points, and when differentiated, also reproduces each component of the gradient at each of the collocation points. Since this technique includes the derivative interpolation explicitly, it eliminates issues with matrix ill-conditioning that can occur in the gradient-enhanced PCE approach based on regression. However, the calculation of high-order global polynomials with the desired interpolation properties can be similarly numerically challenging such that the use of local cubic splines is recommended due to numerical stability.

See Also

These keywords may also be of interest:

- adaptive_sampling
- gpais
- local_reliability
- global_reliability
- sampling
- importance_sampling
- polynomial_chaos

7.2.54.1 p_refinement

- Keywords Area
- method
- stoch_collocation
- p_refinement

Automatic polynomial order refinement
Specification

Alias: none
Argument(s): none
Default: no refinement

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>p-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
</tbody>
</table>

Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the uniform refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
7.2.54.1.1 uniform

- Keywords Area
- method
- stoch_collocation
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.54.1.2 dimension_adaptive

- Keywords Area
- method
- stoch_collocation
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
Required/Optional Description of Dakota Keyword Description
Required (Choose One) Group Dimension Adaptivity Estimation Approach (Group 1) sobol

Dakota Keyword Description

<table>
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<tr>
<th>Required/Optional Description of Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One) Group Dimension Adaptivity Estimation Approach (Group 1)</td>
<td>sobol</td>
</tr>
<tr>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>

### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.54.1.2.1 sobol

- Keywords Area
- method
- stoch_collocation
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized

### Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method,  
  polynomial_chaos  
    sparse_grid_level = 3  
    dimension_adaptive p_refinement sobol  
    max_iterations = 20  
    convergence_tol = 1.e-4
```

**Employ h-refinement to refine the grid**

```plaintext
7.2.54.2 h_refinement
  
  • Keywords Area  
  • method  
  • stoch_collocation  
  • h_refinement
```
### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no refinement

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td></td>
<td>h-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td>dimension_adaptive</td>
<td></td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td></td>
<td>local_adaptive</td>
<td></td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

### Description

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_adaptive options.

#### 7.2.54.2.1 uniform

- Keywords Area
- method
- stoch_collocation
- h_refinement
- uniform

Refine an expansion uniformly in all dimensions.
Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.54.2.2 dimension_adaptive

- Keywords Area
- method
- stoch_collocation
- h_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

Specification

Alias: none
Argument(s): none

Child Keywords:

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
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<td>Required (Choose One)</td>
<td>Dimension Adaptivity Estimation Approach (Group 1)</td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
</tr>
</tbody>
</table>
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.54.2.2.1 sobol

- Keywords Area
- method
- stoch_collocation
- h_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method, polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement generalized
    max_iterations = 20
    convergence_tol = 1.e-4
```

**7.2.54.2.3 local_adaptive**

**Specification**

**Alias:** none

**Argument(s):** none

Planned future capability for local pointwise refinement within a generalized sparse grid.
Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

7.2.54.3  max_refinement_iterations

- Keywords Area
- method
- stoch_collocation
- max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

```
method, polynomial_chaos
dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

7.2.54.4  convergence_tolerance

- Keywords Area
- method
- stoch_collocation
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls
Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See `nl2sol`
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.54.5 `metric_scale`

- Keywords Area
- method
- stoch_collocation
- metric_scale

define scaling of statistical metrics when adapting UQ surrogates

Specification

Alias: none
Argument(s): none
Default: relative
Child Keywords:
### Description

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The **absolute** option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The **relative** option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

The default is **relative**.

**Expected Output**

**Usage Tips**

**Additional Discussion**

#### 7.2.54.5.1 relative

- **Keywords Area**
- **method**
- **stoch_collocation**
- **metric_scale**
- **relative**

relative scaling of statistical metrics when adapting UQ surrogates

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

When adapting a UQ surrogate model, the **relative** option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

**Expected Output**

**Usage Tips**
### Additional Discussion

#### 7.2.54.5.2 absolute

- Keywords Area
- method
- stoch_collocation
- metric_scale
- absolute

Use of absolute statistical metrics when adapting UQ surrogates

### Specification

**Alias:** none  
**Argument(s):** none

### Description

When adapting a UQ surrogate model, the `absolute` option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

### Default Behavior

### Expected Output

### Usage Tips

### Additional Discussion

#### 7.2.54.6 quadrature_order

- Keywords Area
- method
- stoch_collocation
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

### Specification

**Alias:** none  
**Argument(s):** INTEGER

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<th>Optional</th>
<th>Quadrature Rule Nesting (Group 1)</th>
<th>A set of weights specifying the relative importance of each uncertain variable (dimension)</th>
<th>Enforce use of nested quadrature rules if available</th>
</tr>
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<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>nested</td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with `quadrature_order` and, optionally, `dimension_preference`). The default rule selection is to employ `non_nested` Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of `p_refinement` or the case of an explicit `nested` override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a `dimension_preference`, where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified `quadrature_order` and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n}(p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the `quadrature_order` specification supports an array input.

A corresponding sequence specification is documented at, e.g., `quadrature_order_sequence` and `quadrature_order_sequence`.

### See Also

These keywords may also be of interest:

- `quadrature_order_sequence` and `method-multifidelity_stoch_collocation-quadrature_order_sequence`

### 7.2.54.6.1 dimension_preference

- Keywords Area
- method
- stoch_collocation
- quadrature_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.54.6.2 nested

- **Keywords Area**
- method
- stoch\_collocation
- quadrature\_order
- nested

Enforce use of nested quadrature rules if available

Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non\_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.54.6.3 non\_nested

- **Keywords Area**
- method
- stoch\_collocation
- quadrature\_order
- non\_nested

Enforce use of non-nested quadrature rules
7.2 method

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.54.7 sparse_grid_level

- Keywords Area
- method
- stoch_collocation
- sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
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<td>Employ a nodal sparse grid construction in stochastic collocation</td>
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<td>Employ a hierarchical sparse grid construction</td>
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<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 2)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
</tbody>
</table>
Optional (Choose One) | Quadrature Rule Nesting (Group 3) | non_nested |
--- | --- | ---
unrestricted | Enforce use of non-nested quadrature rules if available |

### Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User’s Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User’s Manual for additional details). As for `quadrature_order`, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies. A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`.

### See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence

7.2.54.7.1 `dimension_preference`

- Keywords Area
- method
- stoch_collocation
- sparse_grid_level
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)
7.2 method

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.54.7.2 nodal

- **Keywords Area**
- method
- stoch_collocation
- sparse_grid_level
- nodal

Employ a nodal sparse grid construction in stochastic collocation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** nodal

**Description**

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.

7.2.54.7.3 hierarchical

- **Keywords Area**
- method
- stoch_collocation
- sparse_grid_level
- hierarchical

Employ a hierarchical sparse grid construction
Specification

Alias: none
Argument(s): none

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.54.7.4 restricted

- Keywords Area
- method
- stoch_collocation
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

Alias: none
Argument(s): none
Default: restricted (except for generalized sparse grids)

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of $m = 2l + 1$ are used to enforce odd quadrature orders, where $l$ is the grid level and $m$ is the number of points in the rule. The precision of this Gauss rule is then $i = 2m - 1 = 4l + 1$. For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level $l$ for a sparse grid ($i = 4l + 1$) or an order $m$ for a tensor grid ($i = 2m - 1$). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.54.7.5 unrestricted

- Keywords Area
- method
- stoch_collocation
• **sparse_grid_level**

• **unrestricted**

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

### Specification

**Alias:** none

**Argument(s):** none

### Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

### 7.2.54.7.6 nested

• **Keywords** Area

• **method**

• **stoch_collocation**

• **sparse_grid_level**

• **nested**

Enforce use of nested quadrature rules if available

### Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

### Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are
7.2.54.7.7  non_nested

- Keywords Area
- method
- stoch_collocation
- sparse_grid_level
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.54.8  piecewise

- Keywords Area
- method
- stoch_collocation
- piecewise

Use piecewise local basis functions

Specification

Alias: none
Argument(s): none

Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.
Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)

**Description**

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

**See Also**

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)
Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.54.11 use_derivatives

- Keywords Area
- method
- stoch_collocation
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it’s use with Surfpack Gaussian process is not recommended.

7.2.54.12 samples_on_emulator

- Keywords Area
- method
- stoch_collocation
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples
Argument(s): INTEGER
7.2 method

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
  polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.54.13 sample_type

• Keywords Area

• method

• stoch_collocation

• sample_type

Selection of sampling strategy

Specification

Alias: none

Argument(s): none

Default: random

Child Keywords:

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<th>Description of Group Sample Type (Group 1)</th>
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<th>Dakota Keyword Description</th>
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<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
<td></td>
</tr>
</tbody>
</table>

Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

Default Behavior
If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: `multilevel_sampling` uses Monte Carlo by default).

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

7.2.54.13.1 lhs

- Keywords Area
- method
- stoch_collocation
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: `multilevel_sampling`). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.54.13.2 random

- Keywords Area
- method
• stoch_collocation
• sample_type
• random

Uses purely random Monte Carlo sampling to sample variables

Specification

Alias: none
Argument(s): none

Description

The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```
method sampling
  sample_type random
  samples = 200
```

7.2.54.14 rng

• Keywords Area
• method
• stoch_collocation
• rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none
Default: Mersenne twister (mt19937)
Child Keywords:
### Description

The `rng` keyword is used to indicate a choice of random number generator.

### Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

### Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

### Examples

```plaintext
method
   sampling
      sample_type lhs
      samples = 10
      seed = 98765
      rng rnum2
```

```
7.2.54.14.1 mt19937
```

- Keywords Area
- method
- stoch_collocation
- rng
- mt19937

Generates random numbers using the Mersenne twister

### Specification

**Alias:** none

**Argument(s):** none

### Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).
Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.54.14.2  `rnum2`

- Keywords Area
- `method`
- `stoch_collocation`
- `rng`
- `rnum2`

Generates pseudo-random numbers using the Pecos package

Specification

**Alias:** none

**Argument(s):** none

Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```
7.2.54.15 probability_refinement

- Keywords Area
- method
- stoch_collocation
- probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: sample_refinement

Argument(s): none

Default: no refinement

Child Keywords:

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<td>Importance sampling option for probability refinement</td>
<td>import</td>
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<td>Importance sampling option for probability refinement</td>
<td>adapt_import</td>
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<td>Importance sampling option for probability refinement</td>
<td>mm_adapt_import</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design.</td>
<td></td>
</tr>
</tbody>
</table>

Description

The probability_refinement allows refinement of probability and generalized reliability results using importance sampling. If one specifies probability_refinement, there are some additional options. One can specify which type of importance sampling to use (import, adapt_import, or mm_adapt_import). Additionally, one can specify the number of refinement samples to use with refinement_samples and the seed to use with seed.

The probability_refinement density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.
7.2.54.15.1 import

- Keywords Area
- method
- stoch_collocation
- probability_refinement
- import

Importance sampling option for probability refinement

Specification

**Alias:** none

**Argument(s):** none

Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

7.2.54.15.2 adapt_import

- Keywords Area
- method
- stoch_collocation
- probability_refinement
- adapt_import

Importance sampling option for probability refinement

Specification

**Alias:** none

**Argument(s):** none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.
7.2.54.15.3 mm_adapt_import

- Keywords Area
- method
- stoch_collocation
- probability_refinement
- mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

`mm_adapt_import` starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then `mm_adapt_import` proceeds similarly to `adapt_import` (sample until convergence).

7.2.54.15.4 refinement_samples

- Keywords Area
- method
- stoch_collocation
- probability_refinement
- refinement_samples

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.
Output moments of the specified type and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** standard  

**Child Keywords:**

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<th>Dakota Keyword</th>
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<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

**Description**

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

**Examples**

The following method specification overrides the default to print `central` moments and include them in the set of final statistics.

```plaintext
method, sampling
```
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments central

Overriding the default to none as follows:

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none

allows associated nested model mappings to be simplified from:

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.

7.2.54.16.1 none

• Keywords Area
• method
• stoch_collocation
• final_moments
• none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
7.2 method

7.2.54.16.2 standard
- Keywords Area
- method
- stoch_collocation
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments standard
```

7.2.54.16.3 central
- Keywords Area
- method
- stoch_collocation
- final_moments
- central

Output central moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

**7.2.54.17 response_levels**

- **Keywords Area**
- **method**
- **stoch_collocation**
- **response_levels**

Values at which to estimate desired statistics for each response

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**
If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```bash
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```bash
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n', 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u', 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w', 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h', 'TF2h'
histogram_point_uncertain = 1
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
```
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
<td></td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.4000000000e-05</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>4.0000000000e-05</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>1.0341896485e-05</td>
<td></td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>4.2844660868e-06</td>
<td></td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>8.6000000000e-06</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>1.8000000000e-06</td>
<td></td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.4000000000e-05</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>4.0000000000e-05</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>1.0341896485e-05</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>4.2844660868e-06</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>8.6000000000e-06</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>1.8000000000e-06</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.54.17.1 num_response_levels

- Keywords Area
- method
- stoch_collocation
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.54.17.2 compute

- Keywords Area
- method
- stoch_collocation
- response_levels
- compute

Selection of statistics to compute at each response level
Specification

Alias: none
Argument(s): none
Default: probabilities

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>system</td>
<td>Compute system reliability (series or parallel)</td>
<td></td>
</tr>
</tbody>
</table>

Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```dakota
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```

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7.2 method

3.5e+05 4.0e+05 4.5e+05
compute reliabilities

7.2.54.17.2.1 probabilities

- Keywords Area
- method
- stoch_collocation
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

method sampling
  sample_type random
  samples = 100 seed = 1
  complementary_distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute probabilities

7.2.54.17.2.2 reliabilities

- Keywords Area
- method
- stoch_collocation
- response_levels
The `reliabilities` keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the reliabilities are not computed by default. To change this behavior, the `reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

Computes generalized reliabilities associated with response levels

Specification

**Alias:** none

**Argument(s):** none
Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                 6.0e+04 6.5e+04 7.0e+04
                 3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

Compute system reliability (series or parallel)

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required(Choose One)</strong></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
<td></td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.54.17.2.5 series

- Keywords Area
- method
- stoch_collocation
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.54.17.2.6 parallel

- Keywords Area
- method
- stoch_collocation
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system
7.2.54.18 probability_levels

- Keywords Area
- method
- stoch_collocation
- probability_levels

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Examples**

The Dakota input file below specifies a sampling method with probability levels of interest.
method,
samples = 100 seed = 1
complementary distribution
probability_levels = 1. .66 .33 0.
1. .8 .5 0.
1. .3 .2 0.
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
interface,
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'
responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.1454122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465575e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796373090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.54.18.1 num_probability_levels

- Keywords Area
- method
- stoch_collocation
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.54.19 reliability_levels

- Keywords Area
- method
Specify reliability levels at which the response values will be estimated

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

**7.2.54.19.1 num_reliability_levels**

- **Keywords Area**
- **method**
- **stoch_collocation**
- **reliability_levels**
- **num_reliability_levels**

Specify which reliability_levels correspond to which response

**Specification**

**Alias:** none
7.2 method

**Argument(s):** INTEGERLIST

**Default:** reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.54.20 gen_reliability_levels

- **Keywords Area**
- **method**
- **stoch_collocation**
- **gen_reliability_levels**

Specify generalized reliability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
7.2.54.20.1  num_gen_reliability_levels

- Keywords Area
- method
- stoch_collocation
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.54.21  distribution

- Keywords Area
- method
- stoch_collocation
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
<td></td>
</tr>
</tbody>
</table>
Description

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```dakota
method
    sampling
        sample_type lhs
        samples = 10
        distribution cumulative
```

### 7.2.54.21.1 cumulative

- **Keywords Area**
- **method**
- **stoch_collocation**
- **distribution**
- **cumulative**

Computes statistics according to cumulative functions

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

**Expected Outputs**
Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```
method
   sampling
      sample_type lhs
      samples = 10
      distribution cumulative
```

7.2.54.21.2 complementary

- Keywords Area
- method
- stoch_collocation
- distribution
- complementary

Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```
method
   sampling
      sample_type lhs
      samples = 10
      distribution complementary
```

7.2.54.22 variance_based_decomp

- Keywords Area
Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no variance-based decomposition  

**Child Keywords:**

<table>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>interaction_order</td>
<td></td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
<tr>
<td>Optional</td>
<td>drop_tolerance</td>
<td></td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

**Default Behavior**

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

**Expected Outputs**

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

**Usage Tips**

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.
Examples

```plaintext
method,
  polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of [76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in [90].

7.2.54.22.1 interaction_order

- Keywords Area
- method
- stoch_collocation
- variance_based_decomp
- interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

Description

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.2.54.22.2 drop_tolerance

- Keywords Area
- method
- stoch_collocation
- variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance.
7.2 method

Specification

Alias: none

Argument(s): REAL

Default: All VBD indices displayed

Description

The \texttt{drop\_tolerance} keyword allows the user to specify a value below which sensitivity indices generated by \texttt{variance\_based\_decomp} are not displayed.

Default Behavior

By default, all sensitivity indices generated by \texttt{variance\_based\_decomp} are displayed.

Usage Tips

For \texttt{polynomial\_chaos}, which outputs main, interaction, and total effects by default, the \texttt{univariate\_effects} may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be \texttt{diagonal\_covariance} or \texttt{full\_covariance}, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

```dakota
method,
  sampling
    sample\_type \texttt{lhs}
    samples = 100
  variance\_based\_decomp
    drop\_tolerance = 0.001
```

7.2.54.23 diagonal\_covariance

- Keywords Area
- method
- stoch\_collocation
- diagonal\_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none

Argument(s): none

Default: \texttt{diagonal\_covariance} for response vector $> 10$; else \texttt{full\_covariance}

Description

With a large number of responses, the covariance matrix can be very large. \texttt{diagonal\_covariance} is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
7.2.54.24  full_covariance

- Keywords Area
- method
- stoch_collocation
- full_covariance

Display the full covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.54.25  import_approx_points_file

- Keywords Area
- method
- stoch_collocation
- import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
7.2 method

<table>
<thead>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>freeform</td>
<td></td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

Default Behavior  No import of points at which to evaluate the surrogate.

Expected Output  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

Examples

    method polynomial_chaos
        expansion_order = 4
        import_approx_points_file = 'import.mcmc_annot.dat'
        annotated

7.2.54.25.1  custom_annotated

- Keywords Area
- method
- stoch_collocation
- import_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias:  none

Argument(s):  none

Default:  annotated format

Child Keywords:
## Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
% eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

---

### Keywords Area

- header
- eval_id
- interface_id

- method
Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no header

**Description**

See description of parent custom_annotated

---

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.54.25.2 annotated

• Keywords Area
• method
• stoch_collocation
• import_approx_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.54.25.3 freeform

• Keywords Area

• method

• stoch_collocation

• import_approx_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format
Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
Resulting tabular file:
```

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.54.25.4 active_only

• Keywords Area
  • method
    • stoch_collocation
      • import_approx_points_file
        • active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats
### 7.2 Method

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

---

### 7.2.54.26 export_approx_points_file

- **Keywords Area**
  - method
    - stoch_collocation
    - export_approx_points_file

Output file for surrogate model value evaluations

**Specification**

**Alias:** export_points_file  
**Argument(s):** STRING  
**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

 Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate's predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:
• annotated (default)
  • custom_annotated
  • freeform

7.2.54.26.1 custom_annotated

  • Keywords Area
  • method
  • stoch_collocation
  • export_approx_points_file
  • custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

  • file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.
7.2 Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.54.26.1.1 header

- Keywords Area
- method
- stoch_collocation
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated
7.2.54.26.1.2 eval_id

- Keywords Area
- method
- stoch_collocation
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.54.26.1.3 interface_id

- Keywords Area
- method
- stoch_collocation
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated
7.2.54.26.2 annotated

- Keywords Area
- method
- stoch_collocation
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>@eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.54.26.3 freeform

- Keywords Area
- method
- stoch_collocation
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.54.27  seed

• Keywords Area
  • method
  • stoch_collocation
  • seed

Seed of the random number generator

Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.
Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.54.28 fixed_seed

- Keywords Area
- method
- stoch_collocation
- fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed

7.2.54.29 model_pointer

- Keywords Area
- method
- stoch_collocation
- model_pointer

Identifier for model block to be used by a method
Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
  tabular_data
    method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'
```
variables
uniform_uncertain = 2
lower_bounds = 0.0
upper_bounds = 1.1
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.55 multifidelity_stoch_collocation

• Keywords Area

• method

• multifidelity_stoch_collocation

Multifidelity uncertainty quantification using stochastic collocation

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Automated Refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>h_refinement</td>
<td>Employ h-refinement to refine the grid</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Option</td>
<td>Setting</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>--------------</td>
<td>------------------</td>
<td>------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>metric_scale</td>
<td>define scaling of statistical metrics when adapting UQ surrogates</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>statistics_mode</td>
<td>type of statistical metric roll-up for multifidelity UQ methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
<td></td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Interpolation Grid Type (Group 2)</td>
<td>quadrature_order_sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>sparse_grid_level_sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td>Optional</td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_refinement</td>
<td>Allow refinement of probability and generalized reliability results using importance sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-------------------</td>
<td>-------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>(Choose One) Covariance Type (Group 4) diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_approx_points_file</td>
<td>Filename for points at which to evaluate the PCE/SC surrogate</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td>Sequence of seed values for multi-stage random sampling</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

As described in stoch_collocation, stochastic collocation is a general framework for approximate representation of random response functions in terms of finite-dimensional interpolation bases, using interpolation polynomials that...
may be either local or global and either value-based or gradient-enhanced.

In the multifidelity case, we decompose this interpolant expansion into several constituent expansions, one per model form or solution control level. In a bi-fidelity case with low-fidelity (LF) and high-fidelity (HF) models and an additive discrepancy approach, we have:

\[ R = \sum_{i=1}^{N_{LF}} r_{i}^{LF} L_i(\xi) + \sum_{i=1}^{N_{HF}} \delta_i L_i(\xi) \]

where \( \delta_i \) is a coefficient for the discrepancy expansion.

The same specification options are available as described in stoch_collocation with one key difference: the coefficient estimation inputs change from a scalar input for a single expansion to a sequence specification for a low-fidelity expansion followed by multiple discrepancy expansions.

To obtain the coefficients \( r_i \) and \( \delta_i \) for each of the expansions, the following options are provided:

1. multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order_sequence, and, optionally, dimension_preference).
2. multidimensional integration by the Smolyak sparse grid method (specified with sparse_grid_level_sequence and, optionally, dimension_preference)

It is important to note that, while quadrature_order_sequence and sparse_grid_level_sequence are array inputs, only one scalar from these arrays is active at a time for a particular expansion estimation. In order to specify anisotropy in resolution across the random variable set, a dimension_preference specification can be used to augment these scalar specifications.

Multifidelity UQ using SC requires that the model selected for iteration by the method specification is a multifidelity surrogate model (see hierarchical), which defines an ordered_model_sequence (see hierarchical). Two types of hierarchies are supported: (i) a hierarchy of model forms composed from more than one model within the ordered_model_sequence, or (ii) a hierarchy of discretization levels comprised from a single model within the ordered_model_sequence which in turn specifies a solution_level_control (see solution_level_control).

In both cases, an expansion will first be formed for the low fidelity model or coarse discretization, using the first value within the coefficient estimation sequence, along with any specified refinement strategy. Second, expansions are formed for one or more model discrepancies (the difference between response results if additive correction or the ratio of results if multiplicative correction), using all subsequent values in the coefficient estimation sequence (if the sequence does not provide a new value, then the previous value is reused) along with any specified refinement strategy. The number of discrepancy expansions is determined by the number of model forms or discretization levels in the hierarchy.

After formation and refinement of the constituent expansions, each of the expansions is combined (added or multiplied) into an expansion that approximates the high fidelity model, from which the final set of statistics are generated.

Additional Resources

Dakota provides access to multifidelity SC methods through the NonDMultilevelStochCollocation class. Refer to the Stochastic Expansion Methods chapter of the Theory Manual[16] for additional information on the Multifidelity SC algorithm.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Integration and Expansion Moments (expansion moments only)
- Probability Density
- Level Mappings

In addition, the execution group has the attribute equiv_hf_evals, which records the equivalent number of high-fidelity evaluations.
Examples

```plaintext
method,
    multifidelity_stoch_collocation
    model_pointer = 'HIERARCH'
    sparse_grid_level_sequence = 4 3 2

model,
    id_model = 'HIERARCH'
    surrogate hierarchical
    ordered_model_fidelities = 'LF' 'MF' 'HF'
    correction additive zeroth_order
```

See Also

These keywords may also be of interest:

- `adaptive_sampling`
- `gpais`
- `local_reliability`
- `global_reliability`
- `sampling`
- `importance_sampling`
- `stoch_collocation`

### 7.2.55.1 \texttt{p_refinement}

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **\texttt{p_refinement}**

Automatic polynomial order refinement

**Specification**

- **Alias:** none
- **Argument(s):** none
- **Default:** no refinement
- **Child Keywords:**
### Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

#### 7.2.55.1 uniform

- Keywords Area
- method
- `multifidelity_stoch_collocation`
- `p_refinement`
- `uniform`
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.55.1.2 dimension_adaptive

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **p_refinement**
- **dimension_adaptive**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Dimension Adaptivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol's sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>
Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher 'importance' in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.55.1.2.1 sobol

- Keywords Area
- method
- multifidelity_stoch_collocation
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4

7.2.55.1.2.2 generalized

- Keywords Area
- method
- multifidelity_stoch_collocation
- p_refinement
- dimension_adaptive
- generalized
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method, polynomial_chaos
   sparse_grid_level = 3
   dimension_adaptive p_refinement generalized
   max_iterations = 20
   convergence_tol = 1.e-4
```

7.2.55.2 h_refinement

- **Keywords Area**
  - method
  - multifidelity_stoch_collocation
  - h_refinement

Employ h-refinement to refine the grid.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no refinement

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>h-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
</tbody>
</table>
Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

Planned future capability for local pointwise refinement within a generalized sparse grid.

Description

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_adaptive options.

7.2.55.2.1 uniform

- Keywords Area
- method
- multifidelity_stoch_collocation
- h_refinement
- uniform

Refine an expansion uniformly in all dimensions.

Specification

Alias: none

Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.
Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Dimension Adaptivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

**7.2.55.2.2 sobol**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **h_refinement**
Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** generalized

**Description**

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

**Examples**

```
method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.55.2.2.2 **generalized**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **h_refinement**
- **dimension_adaptive**
- **generalized**

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).
7.2 method

Examples

method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.55.2.3 local_adaptive

  • Keywords Area
  • method
  • multifidelity_stoch_collocation
  • h_refinement
  • local_adaptive

Planned future capability for local pointwise refinement within a generalized sparse grid.

Specification

Alias: none
Argument(s): none

Description

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

7.2.55.3 max_refinement_iterations

  • Keywords Area
  • method
  • multifidelity_stoch_collocation
  • max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

method,
  polynomial_chaos
dimension_adaptive generalized
  p_refinement
    max_refinement_iterations = 20
    convergence_tol = 1.e-4
  sparse_grid_level = 1

7.2.55.4 convergence_tolerance

  • Keywords Area
  • method
  • multifidelity_stoch_collocation
  • convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

  • method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

  • DOT: relative tolerance that must be satisfied for two consecutive iterations
  • NL2SOL: See nl2sol
• NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
• NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.55.5 metric_scale

- Keywords Area
- method
- multifidelity_stoch_collocation
- metric_scale

define scaling of statistical metrics when adapting UQ surrogates

Specification

**Alias:** none

**Argument(s):** none

**Default:** relative

**Child Keywords:**

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<tr>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>relative</td>
<td>relative scaling of statistical metrics when adapting UQ surrogates</td>
</tr>
</tbody>
</table>

**Description**

When adapting a UQ surrogate model, e.g., by change in response covariance matrix or level mappings, this option controls the use of scaling. The **absolute** option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance. The **relative** option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

The default is **relative**.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.55.5.1 relative

- Keywords Area
- method
relative scaling of statistical metrics when adapting UQ surrogates

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

When adapting a UQ surrogate model, the `relative` option indicates that the change is measured relative to the current reference value (which evolves during the refinement), such that the convergence tolerance controls the relative change.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.55.5.2  `absolute`

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **metric_scale**
- **absolute**

use of absolute statistical metrics when adapting UQ surrogates

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

When adapting a UQ surrogate model, the `absolute` option indicates that there is no scaling such that the statistical change is compared directly to the convergence tolerance.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**
7.2.55.6  statistics_mode

- Keywords Area
- method
- multifidelity_stoch_collocation
- statistics_mode

type of statistical metric roll-up for multifidelity UQ methods

Specification

**Alias:** none

**Argument(s):** none

**Default:** DEFAULT_EXPANSION_STATS

**Child Keywords:**

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<th>Dakota Keyword</th>
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<tr>
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<td></td>
<td>active</td>
<td>use statistics for the active expansion within multifidelity refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>combined</td>
<td>use statistics for the combined expansion within multifidelity refinement</td>
</tr>
</tbody>
</table>

**Description**

When performing adaptive refinement in multifidelity UQ methods, one can adapt the expansion for a model level/fidelity based on its local statistics (active) or based on the statistics for the aggregate expansion (combined). The latter case may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

**Default Behavior**

Default for multifidelity methods is **combined statistics**.

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.55.6.1  active

- Keywords Area
- method
- multifidelity_stoch_collocation
- statistics_mode
**Keywords Area**

- **active**

use statistics for the active expansion within multifidelity refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the active model level/fidelity. This approach avoids some computational overhead in some cases by avoiding expansion roll-up.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**

7.2.55.6.2 combined

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **statistics_mode**
- **combined**

use statistics for the combined expansion within multifidelity refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

When performing adaptive refinement in multifidelity UQ methods, this option adapts based on the statistics for the aggregate multifidelity expansion. This approach may incur additional computational overhead, but ensures that the refinement occurs in the aggregate multifidelity context.

**Default Behavior**

**Expected Output**

**Usage Tips**

**Additional Discussion**
Sample allocation approach for multifidelity expansions

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
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<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>Multifidelity Sample Allocation Control (Group 1)</td>
<td>greedy</td>
<td>Sample allocation based on greedy refinement within multifidelity stochastic collocation</td>
</tr>
</tbody>
</table>

**Description**

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

**Default Behavior**

The default, when allocation_control is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

**7.2.55.7.1 greedy**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **allocation_control**
- **greedy**

Sample allocation based on greedy refinement within multifidelity stochastic collocation

**Specification**

**Alias:** none  
**Argument(s):** none
**Description**

Multifidelity stochastic collocation supports greedy refinement strategies using tensor and sparse grids for both nodal and hierarchical collocation approaches. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within an integrated competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.

**Examples**

The following example of greedy multifidelity stochastic collocation using nodal interpolation starts from a zeroth-order expansion (a constant) for each level, and generates uniform candidate refinements for each level that are competed in a greedy competition. The number of new samples for the incremented candidate expansion order is determined from the quadrature rules of the new sparse grid level. In this case, the number of candidates for each level is limited to one uniform refinement of the current sparse grid level.

```plaintext
method,
  model_pointer = 'HIERARCH'
  multifidelity_stoch_collocation
  nodal
  allocation_control greedy
  p_refinement uniform
  sparse_grid_level_sequence = 0 unrestricted
  convergence_tolerance 1.e-3
```

The next example employs generalized sparse grids and hierarchical interpolation. Each level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all levels is competed within a unified greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each level and the number of candidates grows rapidly with random dimension and grid level.

```plaintext
method,
  model_pointer = 'HIERARCH'
  multifidelity_stoch_collocation
  hierarchical
  allocation_control greedy
  p_refinement dimension_adaptive generalized
  sparse_grid_level_sequence = 0 unrestricted
  convergence_tolerance 1.e-8
```

**7.2.55.8 discrepancy_emulation**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **discrepancy_emulation**

Formulation for emulation of model discrepancies.

**Specification**

- **Alias**: none
- **Argument(s)**: none
Default: distinct

Child Keywords:
Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.

Examples

```dakota
method, multilevel_polynomial_chaos
  expansion_order_sequence = 2
  collocation_ratio = .9
  orthogonal_matching_pursuit
  discrepancy_emulation recursive
```

7.2.55.8.1 distinct

- Keywords Area
- method
- multifidelity_stoch_collocation
- discrepancy_emulation
- distinct

Distinct formulation for emulation of model discrepancies.

Specification

**Alias**: paired

**Argument(s)**: none
Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.55.8.2 recursive

- Keywords Area
- method
- multifidelity_stoch_collocation
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.55.9 quadrature_order_sequence

- Keywords Area
- method
- multifidelity_stoch_collocation
- quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST
Child Keywords:
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<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

**Description**

Each level entry of the `quadrature_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `quadrature_order`

**See Also**

These keywords may also be of interest:

- `quadrature_order`

**7.2.55.9.1 dimension_preference**

- Keywords Area
- method
- `multifidelity_stoch_collocation`
- `quadrature_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

*Alias*: none

*Argument(s)*: REALLIST

*Default*: isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.55.9.2 nested

Enforce use of nested quadrature rules if available

Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.55.9.3 non_nested

Enforce use of non-nested quadrature rules

Specification

**Alias:** none

**Argument(s):** none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule
7.2.55.10  sparse_grid_level_sequence

• Keywords Area
  • method
  • multifidelity_stoch_collocation
  • sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
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<tr>
<td>Optional (Choose One)</td>
<td>Group 1</td>
<td>nodal</td>
<td>Employ a nodal sparse grid construction</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 2)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 3)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>
Description

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

See Also

These keywords may also be of interest:

- `sparse_grid_level`

### 7.2.55.10.1 dimension_preference

- **Keywords Area**
- **method**
- `multifidelity_stoch_collocation`
- `sparse_grid_level_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- `sobol`
- `decay`

### 7.2.55.10.2 nodal

- **Keywords Area**
- **method**
- `multifidelity_stoch_collocation`
Emploee a nodal sparse grid construction in stochastic collocation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** nodal

**Description**

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.

7.2.55.10.3  hierarchical

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **sparse_grid_level_sequence**
- **hierarchical**

Employ a hierarchical sparse grid construction

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.55.10.4  restricted

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **sparse_grid_level_sequence**
7.2 method

• restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

Alias: none

Argument(s): none

Default: restricted (except for generalized sparse grids)

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.55.10.5 unrestricted

• Keywords Area
  • method
  • multifidelity_stoch_collocation
  • sparse_grid_level_sequence
  • unrestricted

Overide the default restriction of growth rates for nested and non-nested rules that are by defualt synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences. To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.55.10.6 nested

- Keywords Area
- method
- multifidelity_stoch_collocation
- sparse_grid_level_sequence
- nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.55.10.7 non_nested

- Keywords Area
- method
- multifidelity_stoch_collocation
- sparse_grid_level_sequence
- non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule
7.2.55.11 piecewise

- Keywords Area
- method
- multifidelity_stoch_collocation
- piecewise

Use piecewise local basis functions

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)

**Description**

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.55.12 askey

- Keywords Area
- method
- multifidelity_stoch_collocation
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)
Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

7.2.55.13 wiener

- Keywords Area
- method
- multifidelity_stoch_collocation
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

See Also

These keywords may also be of interest:

- polynomial_chaos
- askey
7.2.55.14  use_derivatives

- Keywords Area
- method
- multifidelity_stoch_collocation
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.55.15  samples_on_emulator

- Keywords Area
- method
- multifidelity_stoch_collocation
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: samples
Argument(s): INTEGER

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.
Examples

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method
polynomial_chaos
  quadrature_order  = 2
  samples_on_emulator = 10000
```

7.2.55.16 sample_type

- Keywords Area
- method
- multifidelity_stoch_collocation
- sample_type

Selection of sampling strategy

Specification

Alias: none

Argument(s): none

Default: random

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

Default Behavior

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

Examples

```plaintext
method
  sampling
    sample_type lhs
```
7.2.55.16.1 lhs

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **sample_type**
- **lhs**

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.55.16.2 random

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **sample_type**
- **random**

Uses purely random Monte Carlo sampling to sample variables
Specification

Alias: none
Argument(s): none

Description

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: `multilevel_sampling`). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```
method sampling
  sample_type random
  samples = 200
```

7.2.55.17  `rng`

- Keywords Area
- method
- `multifidelity_stoch_collocation`
- `rng`

Selection of a random number generator

Specification

Alias: none
Argument(s): none

Default: Mersenne twister (`mt19937`)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>RNG Algorithm (Group 1)</td>
<td><code>mt19937</code></td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>rnum2</code></td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>
Description

The `rng` keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

```dakota
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng rnum2
```

7.2.55.17.1  mt19937

- Keywords Area
- method
- `multifidelity_stoch_collocation`
- `rng`
- `mt19937`

Generates random numbers using the Mersenne twister

Specification

Alias: none
Argument(s): none

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```dakota
method sampling
  sample_type lhs
```
Generates pseudo-random numbers using the Pecos package

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

**Examples**

```yaml
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

**Topics**

This keyword is related to the topics:

- reliability_methods
### Specification

**Alias:** sample_refinement

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Group Importance Sampling Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
<td></td>
</tr>
</tbody>
</table>

| Optional | refinement_samples | Number of samples used to refine a probability estimate or sampling design. |

### Description

The **probability_refinement** allows refinement of probability and generalized reliability results using importance sampling. If one specifies **probability_refinement**, there are some additional options. One can specify which type of importance sampling to use (import, adapt_import, or mm_adapt_import). Additionally, one can specify the number of refinement samples to use with **refinement_samples** and the seed to use with **seed**.

The **probability_refinement** density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

#### 7.2.55.18.1 import

- **Keywords Area**
- method
- multifidelity_stoch_collocation
- probability_refinement
- import

Importance sampling option for probability refinement

### Specification

**Alias:** none

**Argument(s):** none
Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

7.2.55.18.2 adapt_import

- Keywords Area
- method
- multifidelity_stoch_collocation
- probability_refinement
- adapt_import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

7.2.55.18.3 mm_adapt_import

- Keywords Area
- method
- multifidelity_stoch_collocation
- probability_refinement
- mm_adapt_import

Importance sampling option for probability refinement

Specification

Alias: none
Argument(s): none

Description

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more
probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then \texttt{mm\_adapt\_import} proceeds similarly to \texttt{adapt\_import} (sample until convergence).

7.2.55.18.4 \textit{refinement\_samples}

- Keywords Area
- method
- multifidelity\_stoch\_collocation
- probability\_refinement
- refinement\_samples

Number of samples used to refine a probability estimate or sampling design.

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST

**Description**

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using uni-modal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial\_samples in a sampling design.

7.2.55.19 \textit{final\_moments}

- Keywords Area
- method
- multifidelity\_stoch\_collocation
- final\_moments

Output moments of the specified type and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** standard  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

**Examples**

The following method specification overrides the default to print `central` moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```
7.2 method

7.2.55.19.1 none

- Keywords Area
- method
- multifidelity_stoch_collocation
- final_moments
- none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

```
method, sampling
samples = 50 seed = 1234
response_levels = 3.6e+11 1.2e+05 3.5e+05
final_moments none
```

7.2.55.19.2 standard

- Keywords Area
- method
- multifidelity_stoch_collocation
- final_moments
- standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.
Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
    sampling
        samples = 50 seed = 1234
        response_levels = 3.6e+11 1.2e+05 3.5e+05
        final_moments standard
data
```

7.2.55.19.3 central

- Keywords Area
- method
- multifidelity_stoch_collocation
- final_moments
- central

Output central moments and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
    sampling
        samples = 50 seed = 1234
        response_levels = 3.6e+11 1.2e+05 3.5e+05
        final_moments central
data
```

7.2.55.20 response_levels

- Keywords Area
- method
- multifidelity_stoch_collocation
- response_levels

Values at which to estimate desired statistics for each response
Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior

If response_levels are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If response_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The num_response_levels is used to specify which arguments of the response_level correspond to which response.

Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num-
The Dakota input file below specifies a sampling method with response levels of interest.

```
method,
sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
interface,
system async evaluation_concurrency = 5
analysis_driver = 'text_book'
responses,
response_functions = 3
no_gradients
no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

```
Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.64315474e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>2.4350000000e-12</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>1.034186485e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>4.0000000000e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.5000000000e+05</td>
<td>4.284460868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
</tbody>
</table>
```
7.2 method

4.0000000000e+05  4.5000000000e+05  1.8000000000e-06

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.55.20.1  num_response_levels

- Keywords Area
- method
- multifidelity_stoch_collocation
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.
Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

method
sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5

7.2.55.20.2 compute

• Keywords Area
• method
• multifidelity_stoch_collocation
• response_levels
• compute

Selection of statistics to compute at each response level

Specification

Alias: none
Argument(s): none
Default: probabilities
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>probabilities</td>
<td>probabilities</td>
<td>computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>reliabilities</td>
<td>reliabilities</td>
<td>computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>gen_reliabilities</td>
<td>gen_reliabilities</td>
<td>computes generalized reliabilities associated with response levels</td>
</tr>
</tbody>
</table>
### Method

<table>
<thead>
<tr>
<th>Optional</th>
<th>System</th>
<th>Compute system reliability (series or parallel)</th>
</tr>
</thead>
</table>

## Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

### Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

### Expected Output

The type of statistics specified by `compute` will be reported for each response level.

### Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

### Examples

```plaintext
method
    sampling
        sample_type random
        samples = 100 seed = 1
        complementary distribution
            response_levels = 3.6e+11 4.0e+11 4.4e+11
                             6.0e+04 6.5e+04 7.0e+04
                             3.5e+05 4.0e+05 4.5e+05
        compute reliabilities

7.2.55.20.2.1 probabilities

    • Keywords Area
    • method
    • multifidelity_stoch_collocation
    • response_levels
    • compute
    • probabilities

Computes probabilities associated with response levels
```

### Specification

**Alias:** none

**Argument(s):** none
Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```plaintext
method
    sampling
        sample_type random
        samples = 100 seed = 1
    complementary distribution
        response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
        3.5e+05 4.0e+05 4.5e+05
    compute probabilities
```

7.2.55.20.2.2 `reliabilities`

- Keywords Area
- method
- multifidelity_stoch_collocation
- response_levels
- compute
- reliabilities

Computes reliabilities associated with response levels

Specification

**Alias**: none

**Argument(s)**: none

Description

The `reliabilities` keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the reliabilities are not computed by default. To change this behavior, the `reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
Examples

method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
  compute reliabilities

7.2.55.20.2.3 gen_reliabilities
  
  • Keywords Area
  • method
  • multifidelity_stoch_collocation
  • response_levels
  • compute
  • gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none

Argument(s): none

Description

The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
Compute system reliability (series or parallel)

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>System Reliability Mode (Group 1)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

**Description**

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.
7.2.55.20.2.6 parallel

• Keywords Area
• method
• multifidelity_stoch_collocation
• response_levels
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

Description

See parent keyword system for description.

7.2.55.21 probability_levels

• Keywords Area
• method
• multifidelity_stoch_collocation
• probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

Default: No CDF/CCDF response levels to compute

Child Keywords:
Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```plaintext
method, sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
                      1.8 .5 0.
                      1.3 .2 0.
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
```

Dakota Keyword Description

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which <code>probability_levels</code> correspond to which response</td>
</tr>
</tbody>
</table>
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

**Probability Density Function (PDF) histograms for each response function:**

**PDF for response_fn_1:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.138774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

**PDF for response_fn_2:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.983945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

**PDF for response_fn_3:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.302836523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

**Level mappings for each response function:**

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

**7.2.55.21.1 num_probability_levels**

- **Keywords Area**
- **method**
• multifidelity_stoch_collocation
• probability_levels
• num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none
Argument(s): INTEGERLIST
Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.55.22 reliability_levels

• Keywords Area
• method
• multifidelity_stoch_collocation
• reliability_levels

Specify reliability levels at which the response values will be estimated

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.55.22.1 num_reliability_levels

• Keywords Area
• method
• multifidelity_stoch_collocation
• reliability_levels
• num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none
Argument(s): INTEGERLIST
Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.55.23 gen_reliability_levels

• Keywords Area
• method
• multifidelity_stoch_collocation
• gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:
Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.55.23.1  num_gen_reliability_levels

- Keywords Area
- method
- multifidelity_stoch_collocation
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.55.24  distribution

- Keywords Area
- method
Selection of cumulative or complementary cumulative functions

Specification

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>cumulative</td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td>complementary</td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
  distribution cumulative
```

7.2.55.24.1 cumulative

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
• distribution
• cumulative

Computes statistics according to cumulative functions

**Specification**

**Alias:** none
**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```dakota
method	sampling
  sample_type lhs
  samples = 10
  distribution cumulative
```

## 7.2.55.24.2 complementary

• Keywords Area

• method

• multifidelity_stoch_collocation

• distribution

• complementary

Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none
**Argument(s):** none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

```
7.2.55.25  variance_based_decomp
```

- Keywords Area
- method
- multifidelity_stoch_collocation
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into main, interaction, and total effects

Specification

 Alias: none
 Argument(s): none
 Default: no variance-based decomposition

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interaction_order</td>
<td>Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.</td>
</tr>
</tbody>
</table>
Description

Dakota can calculate sensitivity indices through variance-based decomposition using the keyword `variance_based_decomp`. This approach decomposes main, interaction, and total effects in order to identify the most important variables and combinations of variables in contributing to the variance of output quantities of interest.

Default Behavior

Because of processing overhead and output volume, `variance_based_decomp` is inactive by default, unless required for dimension-adaptive refinement using Sobol' indices.

Expected Outputs

When `variance_based_decomp` is specified, sensitivity indices for main effects, total effects, and any interaction effects will be reported. Each of these effects represents the percent contribution to the variance in the model response, where main effects include the aggregated set of univariate terms for each individual variable, interaction effects represent the set of mixed terms (the complement of the univariate set), and total effects represent the complete set of terms (univariate and mixed) that contain each individual variable. The aggregated set of main and interaction sensitivity indices will sum to one, whereas the sum of total effects sensitivity indices will be greater than one due to redundant counting of mixed terms.

Usage Tips

An important consideration is that the number of possible interaction terms grows exponentially with dimension and expansion order. To mitigate this, both in terms of compute time and output volume, possible interaction effects are suppressed whenever no contributions are present due to the particular form of an expansion. In addition, the `interaction_order` and `drop_tolerance` controls can further limit the computational and output requirements.

Examples

```
method,
  polynomial_chaos # or stoch_collocation
  sparse_grid_level = 3
  variance_based_decomp interaction_order = 2
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76] : "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[90].
7.2 method

- variance_based_decomp
- interaction_order

Specify the maximum number of variables allowed in an interaction when reporting interaction metrics.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** Unrestricted (VBD includes all interaction orders present in the expansion)

**Description**

The interaction_order option has been added to allow suppression of higher-order interactions, since the output volume (and memory and compute consumption) of these results could be extensive for high dimensional problems (note: the previous univariate_effects specification is equivalent to interaction_order = 1 in the current specification). Similar to suppression of interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to again save compute and memory resources and reduce output volume).

7.255.25.2 drop_tolerance

- Keywords Area  
- method  
- multifidelity_stoch_collocation  
- variance_based_decomp  
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** All VBD indices displayed

**Description**

The drop_tolerance keyword allows the user to specify a value below which sensitivity indices generated by variance_based_decomp are not displayed.

**Default Behavior**

By default, all sensitivity indices generated by variance_based_decomp are displayed.

**Usage Tips**

For polynomial_chaos, which outputs main, interaction, and total effects by default, the univariate_effects may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
Examples

```
method,
sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
  drop_tolerance = 0.001
```

7.2.55.26 **diagonal_covariance**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **diagonal_covariance**

Display only the diagonal terms of the covariance matrix

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Default**: diagonal_covariance for response vector > 10; else full_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. **diagonal_covariance** is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.55.27 **full_covariance**

- **Keywords Area**
- **method**
- **multifidelity_stoch_collocation**
- **full_covariance**

Display the full covariance matrix

**Specification**

- **Alias**: none
- **Argument(s)**: none

**Description**

With a large number of responses, the covariance matrix can be very large. **full_covariance** is used to force Dakota to output the full covariance matrix.
import_approx_points_file

• Keywords Area
• method
• multifidelity_stoch_collocation
• import_approx_points_file

Filename for points at which to evaluate the PCE/SC surrogate

Specification

Alias: none
Argument(s): STRING
Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td></td>
<td></td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td>freeform</td>
<td></td>
<td></td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

Default Behavior  No import of points at which to evaluate the surrogate.

Expected Output  The PCE/SC surrogate model will be evaluated at the list of points (input variable values) provided in the file and results tabulated and/or statistics computed at them, depending on the method context.

Examples

method
  polynomial_chaos
  expansion_order = 4
  import_approx_points_file = 'import.mcmc_annot.dat'
  annotated

7.2.55.28  custom_annotated

• Keywords Area
• method
• multifidelity_stoch_collocation
• **import_approx_points_file**

• **custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

• **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To control which header row and columns are in the input/output, specify **custom.annotated**, followed by options, in the relevant export/import context.

**Usage Tips**

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to **annotated format**, though **freeform** remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.55.28.1.1  header

• Keywords Area
  • method
    • multifidelity_stoch_collocation
    • import_approx_points_file
    • custom_annotated
      • header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated

7.2.55.28.1.2  eval_id

• Keywords Area
  • method
    • multifidelity_stoch_collocation
    • import_approx_points_file
    • custom_annotated
      • eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.55.28.1.3 interface_id

- Keywords Area
- method
- multifidelity_stoch_collocation
- import_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.55.28.2 annotated

- Keywords Area
- method
- multifidelity_stoch_collocation
- import_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
7.2 specifications

**Alias:** none

**Argument(s):** none

**Default:** annotated format

### Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

### Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

### Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

### Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.55.28.3 freeform

- **Keywords Area**
  - method
  - multifidelity_stoch_collocation
  - import_approx_points_file
  - freeform

Selects freeform file format
Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```dakota
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
 0.9  1.1  0.0002  0.26  0.76
 0.90009  1.1  0.0001996404857  0.2601620081  0.759955
 0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.55.28.4 active_only

• Keywords Area
7.2 method

- method
- multifidelity_stoch_collocation
- import_approx_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.55.29 export_approx_points_file

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:
Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.55.29.1 custom_annotated

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
  tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id    x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1   0.9   1.1  0.0002  0.26  0.76
2   0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3   0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
7.2.55.29.1.1 header

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.55.29.1.2 eval_id

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated
7.2.55.29.1.3 interface_id

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent [custom_annotated](#)

7.2.55.29.2 annotated

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9  1.1  0.0002  0.26  0.76
2 NO_ID 0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3 NO_ID 0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.55.29.3 freeform

- Keywords Area
- method
- multifidelity_stoch_collocation
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9      1.1      0.0002    0.26      0.76
0.90009  1.1  0.0001996404857  0.2601620081   0.759955
0.89991  1.1  0.0002003604863  0.2598380081   0.760045
...```

7.2.55.30 seed_sequence

- Keywords Area

- method

- multifidelity_stoch_collocation

- seed_sequence

Sequence of seed values for multi-stage random sampling
Specification

**Alias:** none

**Argument(s):** INTGELIST

**Default:** system-generated (non-repeatable)

**Description**

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

**Default Behavior**

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

**Usage Tips**

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

**Examples**

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
   model_pointer = ’HIERARCH’
   multilevel_sampling
   pilot_samples = 100 50 40 30 20
   seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

**See Also**

These keywords may also be of interest:

- **seed**
- **fixed_seed**

7.255.31 fixed_seed

- **Keywords Area**
Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run

**Description**

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

**Topics**

This keyword is related to the topics:

- `block_pointer`
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                    0.1 0.2 0.6
                    0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0.  0.
upper_bounds = 1.  1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system async evaluation_concurrency = 5
```
7.2 method

```
analysis_driver = 'text_book'
```

responses
response_functions = 3
no_gradients
no_hessians

7.2.56 sampling

- Keywords Area
- method
- sampling

Randomly samples variables according to their distributions

Topics

This keyword is related to the topics:

- uncertainty_quantification
- sampling

Specification

**Alias:** nond_sampling

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>samples</td>
<td>samples</td>
<td>Number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_-samples</td>
<td>refinement_-samples</td>
<td>Performs an incremental Latin Hypercube Sampling (LHS) study</td>
</tr>
<tr>
<td>Optional</td>
<td>d_optimal</td>
<td>Generate a D-optimal sampling design</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-----------</td>
<td>-------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into contributions from variables</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>backfill</td>
<td>Ensures that the samples of discrete variables with finite support are unique</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>principal_components</td>
<td>Activates principal components analysis of the response matrix of N samples * L responses.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>wilks</td>
<td>Number of samples for random sampling using Wilks statistics</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
<td></td>
</tr>
</tbody>
</table>
This method generates parameter values by drawing samples from the specified uncertain variable probability distributions. The computational model is executed over all generated parameter values to compute the responses for which statistics are computed. The statistics support sensitivity analysis and uncertainty quantification.

**Default Behavior**

By default, sampling methods operate on aleatory and epistemic uncertain variables. The types of variables can be restricted or expanded (to include design or state variables) through use of the `active` keyword in the variables block in the Dakota input file. If continuous design and/or state variables are designated as active, the sampling algorithm will treat them as parameters with uniform probability distributions between their upper and lower bounds. Refer to `variable_support` for additional information on supported variable types, with and without correlation.

The following keywords change how the samples are selected:

- `sample_type`
- `fixed_seed`
- `rng`
- `samples`
- `seed`
- `variance_based_decomp`

**Expected Outputs**

As a default, Dakota provides correlation analyses when running LHS. Correlation tables are printed with the simple, partial, and rank correlations between inputs and outputs. These can be useful to get a quick sense of how correlated the inputs are to each other, and how correlated various outputs are to inputs. `variance_based_decomp` is used to request more sensitivity information, with additional cost.

Additional statistics can be computed from the samples using the following keywords:

- `response_levels`
- `reliability_levels`
- `probability_levels`
response_levels computes statistics at the specified response value. The other three allow the specification of the statistic value, and will estimate the corresponding response value.

distribution is used to specify whether the statistic values are from cumulative or complementary cumulative functions.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- When variance_based_decomp is enabled
  - Variance-Based Decomposition (Sobol' Indices)
- For aleatory UQ studies
  - Probability Density
  - Level Mappings
  - Sampling Moments
  - Correlations
- For epistemic UQ studies
  - Extreme Responses
  - Correlations

Usage Tips

sampling is a robust approach to doing sensitivity analysis and uncertainty quantification that can be applied to any problem. It requires more simulations than newer, advanced methods. Thus, an alternative may be preferable if the simulation is computationally expensive.

Examples

# tested on Dakota 6.0 on 140501

environment
tabular_data
tabular_data_file = 'Sampling_basic.dat'

method
sampling
  sample_type lhs
  samples = 20

model
single

variables
active uncertain
uniform_uncertain = 2
  descriptors = 'input1' 'input2'
  lower_bounds = -2.0 -2.0
  upper_bounds = 2.0 2.0
  continuous_state = 1
  descriptors = 'constant1'
  initial_state = 100

interface
analysis_drivers 'text_book'
  fork

responses
This example illustrates a basic sampling Dakota input file.

- LHS is used instead of purely random sampling.
- The default random number generator is used.
- Without a seed specified, this will not be reproducible
- In the variables block, two types of variables are used
- Only the uncertain variables are varied, this is the default behavior, and is also specified by the active keyword, w/ the uncertain option

See Also

These keywords may also be of interest:

- active

FAQ

Q: Do I need to keep the LHS* and S4 files? A: No

7.2.56.1 samples

- Keywords Area
- method
- sampling
- samples

Number of samples for sampling-based methods

Specification

Alias: initial_samples
Argument(s): INTEGER
Default: 0

Description

The samples keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips
To obtain linear sensitivities or to construct a linear response surface, at least \( \text{dim}+1 \) samples should be used, where \( \text{"dim"} \) is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \( \frac{(\text{dim}+1)(\text{dim}+2)}{2} \) samples are needed. For uncertainty quantification, we recommend at least \( 10^\ast \text{dim} \) samples. For \text{variance}_\text{based}_\text{decomp}, we recommend hundreds to thousands of samples. Note that for \text{variance}_\text{based}_\text{decomp}, the number of simulations performed will be \( N^\ast(\text{dim}+2) \).

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20

7.2.56.2   seed

  • Keywords Area
  • method
  • sampling
  • seed

Seed of the random number generator

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

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7.2 method

7.2.56.3 fixed_seed

- Keywords Area
- method
- sampling
- fixed_seed

Reuses the same seed value for multiple random sampling sets

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

7.2.56.4 sample_type

- Keywords Area
- method
- sampling
- sample_type

Selection of sampling strategy

Specification

Alias: none
Argument(s): none
Default: lhs

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required <em>(Choose One)</em></td>
<td>Sample Type (Group 1)</td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>incremental_lhs</td>
<td>(Deprecated keyword) Augments an existing Latin Hypercube Sampling (LHS) study</td>
</tr>
<tr>
<td></td>
<td></td>
<td>incremental_random</td>
<td>(Deprecated keyword) Augments an existing random sampling study</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: `multilevel_sampling` uses Monte Carlo by default).

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

7.2.56.4.1 `lhs`

- Keywords Area
- method
- sampling
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables
### 7.2 method

#### Specification

**Alias:** none  
**Argument(s):** none

#### Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

#### Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

#### Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

#### Examples

```dakota
method
sampling
  sample_type lhs
  samples = 20
```

**7.2.56.4.2 random**

- **Keywords Area**
- **method**
- **sampling**
- **sample_type**
- **random**

Uses purely random Monte Carlo sampling to sample variables

#### Specification

**Alias:** none  
**Argument(s):** none

#### Description

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

#### Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.
Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```
method
  sampling
    sample_type random
    samples = 200
```

7.2.56.4.3 incremental_lhs

- Keywords Area
- method
- sampling
- sample_type
- incremental_lhs

(Deprecated keyword) Augments an existing Latin Hypercube Sampling (LHS) study

Specification

**Alias:** none

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

Description

This keyword is deprecated. Instead specify `sample_type lhs` with `refinement_samples`.

7.2.56.4.4 incremental_random

- Keywords Area
- method
- sampling
- sample_type
- incremental_random

(Deprecated keyword) Augments an existing random sampling study

Specification

**Alias:** none

**Argument(s):** none
7.2.56.5 refinement_samples

- Keywords Area
- method
- sampling
- refinement_samples

Performs an incremental Latin Hypercube Sampling (LHS) study

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST

**Description**

Use of refinement_samples replaces the deprecated sample_type incremental_lhs and sample_type incremental_random.

An incremental random sampling approach will successively add samples to an initial or existing random sampling study according to the sequence of refinement_samples. Dakota reports statistics (mean, variance, percentiles, etc) separately for the initial samples and for each refinement_samples increment at the end of the study. For an LHS design, the number of refinement_samples in each refinement level must result in twice the number of previous samples. For sample_type random, there is no constraint on the number of samples that can be added.

Often, this approach is used when you have an initial study with sample size N1 and you want to perform an additional N1 samples. The initial N1 samples may be contained in a Dakota restart file so only N1 (instead of 2 x N1) potentially expensive function evaluations will be performed.

This process can be extended by repeatedly increasing (for LHS: doubling) the refinement_samples:

```
method
  sampling
    seed = 1337
    samples = 50
    refinement_samples = 50 100 200 400 800
```

**Usage Tips**

The incremental approach is useful if it is uncertain how many simulations can be completed within available time.

See the examples below and the Usage and Restarting Dakota Studies pages.

**Examples**

Suppose an initial study is conducted using sample_type lhs with samples = 50. A follow-on study uses a new input file where samples = 50, and refinement_samples = 50, resulting in 50 reused samples (from restart) and 50 new random samples. The 50 new samples will be combined with the 50 previous samples to generate a combined sample of size 100 for the analysis.

One way to ensure the restart file is saved is to specify a non-default name, via a command line option:
dakota -input LHS_50.in -write_restart LHS_50.rst

which uses the input file:

```
# LHS_50.in
environment
  tabular_data
    tabular_data_file = 'LHS_50.dat'
method
  sampling
    seed = 1337
    sample_type lhs
    samples = 50
model
  single
variables
  uniform_uncertain = 2
  descriptors = 'input1' 'input2'
  lower_bounds = -2.0 -2.0
  upper_bounds = 2.0 2.0
interface
  analysis_drivers 'text_book'
  fork
responses
  response_functions = 1
  no_gradients
  no_hessians

and the restart file is written to LHS_50.rst.

Then an incremental LHS study can be run with:

dakota -input LHS_100.in -read_restart LHS_50.rst -write_restart LHS_100.rst

where LHS_100.in is shown below, and LHS_50.rst is the restart file containing the results of the previous LHS study. In the example input files for the initial and incremental studies, the values for seed match. This ensures that the initial 50 samples generated in both runs are the same.

```
# LHS_100.in
environment
  tabular_data
    tabular_data_file = 'LHS_100.dat'
method
  sampling
    seed = 1337
    sample_type lhs
    samples = 50
    refinement_samples = 50
model
  single
variables
  uniform_uncertain = 2
  descriptors = 'input1' 'input2'
  lower_bounds = -2.0 -2.0
  upper_bounds = 2.0 2.0
interface
  analysis_drivers 'text_book'
  fork
responses
```
The user will get 50 new LHS samples which maintain both the correlation and stratification of the original LHS sample. The new samples will be combined with the original samples to generate a combined sample of size 100.

### 7.2.56.6 d_optimal

- **Keywords Area**
- **method**
- **sampling**
- **d_optimal**

Generate a D-optimal sampling design

#### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** off  

#### Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group Design Strategy (Group 1)</th>
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</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>candidate_designs</td>
<td>Number of candidate sampling designs from which to select the most D-optimal</td>
</tr>
</tbody>
</table>

**Description**

This option will generate a sampling design that is approximately determinant-optimal (D-optimal) by downselecting from a set of candidate sample points.

**Default Behavior**

If not specified, a standard sampling design (MC or LHS) will be generated. When d_optimal is specified, 100 candidate designs will be generated and the most D-optimal will be selected.

**Usage Tips**

D-optimal designs are only supported for aleatory_uncertain_variables. The default candidate-based D-optimal strategy works for all submethods except incremental LHS (lhs with refinement_samples). The Leja sampling option only works for continuous variables, and when used with LHS designs, the candidates point set will be Latin, but the final design will not be.
Examples

```dakota
method
  sampling
    sample_type random
    samples = 20
    d_optimal
  candidate_designs = 500
```

7.2.56.6.1 candidate_designs

- Keywords Area
- method
- sampling
- d_optimal
- candidate_designs

Number of candidate sampling designs from which to select the most D-optimal

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

Description

Dakota will **generate** candidate_designs using the specified sampling strategy and select the one that is more determinant-optimal (D-optimal)

Examples

```dakota
method
  sampling
    sample_type random
    samples = 20
    d_optimal
    candidate_designs = 500
```

7.2.56.6.2 leja_oversample_ratio

- Keywords Area
- method
- sampling
- d_optimal
- leja_oversample_ratio

Oversampling ratio for generating candidate point set
**7.2 method**

**Specification**

Alias: none  
Argument(s): REAL  
Default: 10.0  

**Description**

When generating a D-Optimal point set of size N, the `oversample_ratio` R controls the number of candidate points R x N from which the D-Optimal points are chosen.

**Default Behavior**

The default when not specified is to perform candidate-based sample design selection (non-Leja)

**Examples**

```plaintext
method  
sampling  
  sample_type random  
  samples = 20  
  d_optimal  
    oversample_ratio = 2.0
```

**7.2.56.7 variance_based_decomp**

- **Keywords Area**
- **method**
- **sampling**
- **variance_based_decomp**

Activates global sensitivity analysis based on decomposition of response variance into contributions from variables

**Specification**

Alias: none  
Argument(s): none  
Default: no variance-based decomposition  
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>drop_tolerance</td>
<td></td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>
Description

Dakota can calculate sensitivity indices through variance based decomposition using the keyword `variance_based_decomp`. These indicate how important the uncertainty in each input variable is in contributing to the output variance.

Default Behavior

Because of the computational cost, `variance_based_decomp` is turned off as a default.

If the user specified a number of samples, N, and a number of nondeterministic variables, M, variance-based decomposition requires the evaluation of \( N \times (M+2) \) samples. **Note that specifying this keyword will increase the number of function evaluations above the number requested with the `samples` keyword since replicated sets of sample values are evaluated.**

Expected Outputs

When `variance_based_decomp` is specified, sensitivity indices for main effects and total effects will be reported. Main effects (roughly) represent the percent contribution of each individual variable to the variance in the model response. Total effects represent the percent contribution of each individual variable in combination with all other variables to the variance in the model response.

Usage Tips

To obtain sensitivity indices that are reasonably accurate, we recommend that N, the number of samples, be at least one hundred and preferably several hundred or thousands.

Examples

```plaintext
method,
  sampling
    sample_type lhs
    samples = 100
    variance_based_decomp
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of[76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, \( S_i \), means that the uncertainty in the input variable \( i \) has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in[76] and[90].

7.2.56.7.1 drop_tolerance

- **Keywords Area**
- **method**
- **sampling**
- **variance_based_decomp**
- **drop_tolerance**

Suppresses output of sensitivity indices with values lower than this tolerance.

Specification

**Alias:** none
**7.2 method**

**Argument(s):** REAL  
**Default:** All VBD indices displayed

**Description**

The **drop_tolerance** keyword allows the user to specify a value below which sensitivity indices generated by **variance_based_decomp** are not displayed.

**Default Behavior**

By default, all sensitivity indices generated by **variance_based_decomp** are displayed.

**Usage Tips**

For **polynomial_chaos**, which outputs main, interaction, and total effects by default, the **univariate_effects** may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

**Examples**

```plaintext
method,  
sampling  
  sample_type lhs  
samples = 100  
  variance_based_decomp  
  drop_tolerance = 0.001
```

**7.2.56.8 backfill**

- **Keywords Area**
- **method**
- **sampling**
- **backfill**

Ensures that the samples of discrete variables with finite support are unique

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Traditional LHS can generate replicate samples when applied to discrete variables. This keyword enforces uniqueness, which is determined only over the set of discrete variables with finite support. This allows one to generate LHS for a mixed set of continuous and discrete variables whilst still enforcing that the set of discrete LHS components of all the samples are unique.

**Default Behavior**

Uniqueness of samples over discrete variables is not enforced.

**Usage Tips**
Uniqueness can be useful when applying discrete LHS to simulations without noise.

Examples

```dakota
method,
sampling
    samples = 12
    seed = 123456
    sample_type lhs backfill
variables,
    active all
    uniform_uncertain = 1
    lower_bounds = 0.
    upper_bounds = 1.
    descriptors = 'continuous-uniform'

discrete_uncertain_set
    integer = 1
    elements_per_variable = 4
    elements 1 3 5 7
    descriptors = 'design-set-int'
    real = 1
    initial_point = 0.50
    set_values = 0.25 0.50 0.75 1.00
    descriptors = 'design-set-real'

interface,
    direct analysis_driver = 'text_book'
responses,
    response_functions = 3
    no_gradients
    no_hessians
```

See Also

These keywords may also be of interest:

- `lhs`

7.2.56.9 principal_components

```dakota
    Keywords Area
    method
    sampling
    principal_components
```

Activates principal components analysis of the response matrix of N samples + L responses.

Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required/Optional</td>
<td>percent_variance_explained</td>
<td>Specifies the number of components to retain to explain the specified percent variance.</td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate the principal components of the response matrix of N samples × L responses using the *principal_components* keyword. Principal components analysis (PCA) is a data reduction method. The Dakota implementation is under active development: the PCA capability may ultimately be specified elsewhere or used in different ways. For now, it is performed as a post-processing analysis based on a set of Latin Hypercube samples.

We now have field responses in Dakota. PCA is an initial approach in Dakota to analyze and represent the field data. Specifically, if we have a sample ensemble of field data responses, we want to identify the principal components responsible for the spread of that data. Then, we can generate a surrogate model by representing the overall response as weighted sum of M principal components, where the weights will be determined by GPs which are a function of the input uncertain variables. This reduced form then can be used for sensitivity analysis, calibration, etc.

The steps involved when one specifies *principal_components* in Dakota are as follows:

- Create an LHS input sample based on the uncertain variable specification and run the user-specified model at the LHS points to compute the field responses. For notation purposes, there are d input parameters, N samples, and the field length is L.

- Perform PCA on the covariance matrix of the data set from the previous step. This is done by first centering the data (e.g. subtracting the mean of each column from that column) and performing a singular value decomposition on the covariance matrix of the centered data. The eigenvectors of the covariance matrix correspond to the principal components.

- Identify M principal components based on the percentage of variance explained. There is an optional keyword for *principal_components* called *percent_variance_explained*, which is a threshold that determines the number of components that are retained to explain at least that amount of variance. For example, if the user specifies *percent_variance_explained* = 0.99, the number of components that accounts for at least 99 percent of the variance in the responses will be retained. The default for this percentage is 0.95. In many applications, only a few principal components explain the majority of the variance, resulting in significant data reduction.

- Use the principal components in a predictive sense, by constructing a prediction approximation. The basis functions for this approximation are the principal components. The coefficients of the bases are obtained by constructing GP surrogates for the factor scores of the M principal components. The GP surrogates will be functions of the uncertain inputs. The idea is that we have just performed PCA on (for example) the covariance matrix of 100 samples. Typically, those 100 samples will be generated by sampling over some d uncertain input parameters denoted by u, so there should be a mapping from u to the field data, specifically to the loading coefficients and the factor scores. Currently, the final item printed from a Principal Components Analysis in Dakota is a set of prediction samples based on this prediction approximation or surrogate model that relies on the principal components.

**Default Behavior**

*principal_components* is turned off as a default. It may be used with either scalar responses or field responses, but it is intended to be used with large field responses as a data reduction method. For example, typically we expect the number of LHS samples, N, to be less than the number of field responses, L (e.g. if there is one field, the number of responses values is the length of that field).
**Expected Outputs**

When `principal_components` is specified, the number of significant principal components is printed along with the predictions based on the principal components. If `output debug` is specified, additional information is printed, including the original response matrix, the centered data, the principal components, and the factor scores.

**Usage Tips**

This is a preliminary capability that is undergoing active development. Please contact the Dakota developers team if you have problems with using this capability or want to suggest additional features.

**Examples**

```plaintext
method, sampling
    sample_type lhs
    samples = 100
    principal_components
    percent_variance_explained = 0.98
```

**Theory**

There is an extensive statistical literature available on PCA. We recommend that the interested user peruse some of this in using the PCA capability.

**7.2.56.9.1 percent_variance_explained**

- **Keywords Area**
- **method**
- **sampling**
- **principal_components**
- **percent_variance_explained**

Specifies the number of components to retain to explain the specified percent variance.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Description**

Dakota can calculate the principal components of the response matrix of N samples * L responses using the keyword `principal_components`. Principal components analysis (PCA) is a data reduction method. `percent_variance_explained` is a threshold that determines the number of components that are retained to explain at least that amount of variance. For example, if the user specifies `percent_variance_explained = 0.99`, the number of components that accounts for at least 99 percent of the variance in the responses will be retained. The default for this percentage is 0.95. In many applications, only a few principal components explain the majority of the variance, resulting in significant data reduction.

**Expected Outputs**

**Usage Tips** `percent_variance_explained` should be a real number between 0.0 and 1.0. Typically, it will be between 0.9 and 1.0.
Examples

```
method,
sampling
  sample_type lhs
  samples = 100
  principal_components
  percent_variance_explained = 0.98
```

Theory

There is an extensive statistical literature available on PCA. We recommend that the interested user peruse some of this in using the PCA capability.

7.2.56.10 wilks

- Keywords Area
- method
- sampling
- wilks

Number of samples for random sampling using Wilks statistics

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>order</td>
<td>The order of the statistics to use when determining sample sizes for random sampling using Wilks order statistics.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>confidence_level</td>
<td>The confidence level to be used when determining sample sizes for random sampling using Wilks order statistics.</td>
</tr>
</tbody>
</table>
### Description

The `wilks` keyword is used to compute the number of samples to execute for a random sampling study using Wilks statistics\[91\] and\[68\]. In contrast to most sampling studies where the user specifies the number of samples in advance, Wilks determines the number of samples to run to achieve a particular objective. Specifically, Wilks statistics specify a probability level, \( \alpha \), and confidence level, \( \beta \), and determines the minimum number of samples required such that there is \( \beta \)% confidence that the \( \alpha \times 100 \) percentile of the uncertain distribution on model output will fall below the actual \( \alpha \times 100 \) percentile given by the sample when outputs are ordered from smallest to largest. Statistics can also be either `one_sided` or `two_sided` with the former reflecting a statement about uppermost sample output and the latter reflecting both the smallest and largest sample outputs. Finally, the order of the statistics can be increased to higher order such that the statement concerning probability level and confidence level applies to the uppermost \( M \) outputs (for one-sided \( M \)-order Wilks) or the lowest \( M \) and uppermost \( M \) outputs (for two-sided \( M \)-order Wilks).

### Default Behavior

By default, Wilks statistics are computed using one-sided first-order order statistics with a 95% confidence interval (\( \beta \)) and 95% probability (\( \alpha \)). This results in a sample size of 59.

### Usage Tips

Wilks sample sizes apply to model outputs considered one-at-a-time. Joint variation among multiple outputs requires a generalization of the Wilks approach and is not supported in Dakota at this time.

When more than one probability level is specified, the largest sample size will be performed and used to subsample for the lower probability levels.

### Examples

```dakota
method
   sampling
      sample_type random
```
The order of the statistics to use when determining sample sizes for random sampling using Wilks order statistics.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

**Default Behavior**

The default order is 1.

**Examples**

```
method
  sampling
    sample_type random
  wilks
    order 2
```

**See Also**

These keywords may also be of interest:

- wilks

7.2.56.10.2 confidence_level

The confidence level to be used when determining sample sizes for random sampling using Wilks order statistics.
Specification

**Alias:** none

**Argument(s):** REAL

Description

**Default Behavior**

A value of 0.95 (95%) is the default for both confidence level as well as probability level resulting in 59 samples for first-order one-sided statistics.

Examples

```plaintext
method	sampling
  sample_type random
  wilks
    confidence_level 0.99
```

See Also

These keywords may also be of interest:

- `wilks` method-sampling-probability_levels

7.2.56.10.3 one_sided_lower

- Keywords Area
  - method
    - sampling
  - wilks
    - one_sided_lower

Specifies one-sided lower portion order statistics to be used when determining sample sizes for random sampling using Wilks order statistics.

Specification

**Alias:** none

**Argument(s):** none

Description

This option causes the sample size to be based on the lowermost N outputs for order N statistics (default N=1).

Examples

```plaintext
method	sampling
  sample_type random
  wilks
    one_sided_lower
```
7.2 method

See Also

These keywords may also be of interest:

- wilks

7.2.56.10.4 one_sided_upper

- Keywords Area
- method
- sampling
- wilks
- one_sided_upper

Specifies one-sided upper portion order statistics to be used when determining sample sizes for random sampling using Wilks order statistics.

Specification

Alias: none
Argument(s): none

Description

This option causes the sample size to be based on the uppermost N outputs for order N statistics (default N=1).

Default Behavior  This is the default for Wilks.

Examples

```
method
  sampling
    sample_type random
  wilks
    one_sided_upper
```

See Also

These keywords may also be of interest:

- wilks

7.2.56.10.5 two_sided

- Keywords Area
- method
- sampling
- wilks
- two_sided

Specifies two-sided order statistics (an interval) to be used when determining sample sizes for random sampling using Wilks order statistics.
### Specification

**Alias:** none  
**Argument(s):** none

### Description

This option causes the sample size to be based on the interval defined by the lowest N and uppermost N outputs for order N statistics (default N=1).

**Default Behavior**  The default for Wilks is one_sided_upper.

### Examples

```
method
  sampling
    sample_type random
    wilks
      two_sided
```

### See Also

These keywords may also be of interest:

- wilks

### Output moments of the specified type and include them within the set of final statistics.

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** standard

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
</tbody>
</table>
When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

### Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none
```

takes associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```
Omit moments from the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

**Examples**

```plaintext
method, sampling
  samples = 50  seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

Output standardized moments and include them within the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.
Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, sampling
  samples = 50  seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments standard
```

### 7.2.56.11.3 central

- **Keywords Area**
- **method**
- **sampling**
- **final_moments**
- **central**

Output central moments and include them within the set of final statistics.

### Specification

**Alias:** none

**Argument(s):** none

### Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

### Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method, sampling
  samples = 50  seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

### 7.2.56.12 response_levels

- **Keywords Area**
- **method**
- **sampling**
- **response_levels**

Values at which to estimate desired statistics for each response.
Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior

If response_levels are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If response_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The num_response_levels is used to specify which arguments of the response_level correspond to which response.

Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num-
method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.56.12.1 num_response_levels

- Keywords Area
- method
- sampling
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.
Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```plaintext
method
sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.56.12.2 compute

- **Keywords Area**
  - method
  - sampling
  - response_levels
  - compute

Selection of statistics to compute at each response level

Specification

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>probabilities</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>reliabilities</td>
<td>reliabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td>gen_reliabilities</td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
</tbody>
</table>
The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```
method sampling
  sample_type random
  samples = 100 seed = 1
complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
compute reliabilities
```

**7.2.56.12.2.1 probabilities**

- **Keywords Area**
- **method**
- **sampling**
- **response_levels**
- **compute**
- **probabilities**

Computes probabilities associated with response levels

**Specification**

- **Alias**: none
- **Argument(s)**: none
Description

The probabilities keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the probabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```
method sampling
   sample_type random
   samples = 100 seed = 1
   complementary distribution
   response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
   compute probabilities
```

7.2.56.12.2.2 reliabilities

- Keywords Area
- method
- sampling
- response_levels
- compute
- reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none

Argument(s): none

Description

The reliabilities keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the reliabilities are not computed by default. To change this behavior, the reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.
Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

7.2.56.12.2.3 gen_reliabilities

- Keywords Area
- method
- sampling
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

- **Alias:** none
- **Argument(s):** none

Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```
Compute system reliability (series or parallel)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required</strong>&lt;br&gt;(Choose One)</td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

**Description**

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

**7.2.56.12.2.5 series**

- Keywords Area
- method
- sampling
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system
Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.56.12.2.6 parallel

- Keywords Area
- method
- sampling
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.56.13 probability_levels

- Keywords Area
- method
- sampling
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute
Child Keywords:
Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method, sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
  1. .8 .5 0.
  1. .3 .2 0.
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
```
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.419614379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.160381379e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.160381379e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.419614379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.160381379e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.56.13.1 num_probability_levels

- Keywords Area
- method
7.2 method

- sampling
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.56.14 reliability_levels

- Keywords Area
- method
- sampling
- reliability_levels

Specify reliability levels at which the response values will be estimated

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_reliability_levels</td>
<td></td>
<td>Specify which reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.56.14.1  num_reliability_levels

- Keywords Area
- method
- sampling
- reliability_levels
- num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: reliability_levels evenly distributed among response functions

Description

See parent page

7.2.56.15  gen_reliability_levels

- Keywords Area
- method
- sampling
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.56.15.1 num_gen_reliability_levels

- Keywords Area
- method
- sampling
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: gen_reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.56.16 distribution

- Keywords Area
- method
Keywords Area

- sampling
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
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<tr>
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<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.56.16.1 cumulative

- Keywords Area
- method
- sampling
7.2 method

- distribution
- cumulative

Computes statistics according to cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must appear in conjunction with the distribution keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.56.16.2 complementary

- Keywords Area
- method
- sampling
- distribution
- complementary

Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none
Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

7.2.56.17  **rng**

- **Keywords Area**
- **method**
- **sampling**
- **rng**

Selection of a random number generator

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister ( mt19937 )

**Child Keywords:**

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<th>Dakota Keyword</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.
Default Behavior

If specified, the \texttt{rng} keyword must be accompanied by either \texttt{rnum2} (pseudo-random numbers) or \texttt{mt19937} (random numbers generated by the Mersenne twister). Otherwise, \texttt{mt19937}, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

\begin{verbatim}
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng rnum2
\end{verbatim}

7.2.56.17.1 \texttt{mt19937}

- Keywords Area
- method
- sampling
- rng
- \texttt{mt19937}

Generates random numbers using the Mersenne twister

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: none

Description

The \texttt{mt19937} keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: \url{http://en.wikipedia.org/wiki/Mersenne_twister}.

Default Behavior

\texttt{mt19937} is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended.

Examples

\begin{verbatim}
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng mt19937
\end{verbatim}
### 7.2.56.17.2  rnum2

- Keywords Area
- method
- sampling
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

#### Specification

**Alias:** none  
**Argument(s):** none

#### Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

#### Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

#### Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

#### Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
  rng rnum2
```

### 7.2.56.18  model_pointer

- Keywords Area
- method
- sampling
- model_pointer

Identifier for model block to be used by a method

#### Topics

This keyword is related to the topics:

- block_pointer
7.2 method

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See [block_pointer](#) for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.57 multilevel_sampling

- Keywords Area
- method
- multilevel_sampling

Multilevel sampling methods for UQ

Specification

Alias: multilevel_mc mlmc

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td></td>
<td>Sequence of seed values for multi-stage random sampling</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td></td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>pilot_samples</td>
<td></td>
<td>Initial set of samples for multilevel sampling methods.</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td></td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>export_sample_sequence</td>
<td></td>
<td>Enable export of multi-level/multifidelity sample sequences to individual files</td>
</tr>
<tr>
<td>Optional</td>
<td>allocation_target</td>
<td>Allocation statistics/target for the MLMC sample allocation.</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-------------------</td>
<td>---------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>qoi_aggregation</td>
<td>Aggregation strategy for the QoIs statistics for problems with multiple responses in the MLMC algorithm</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on relative error</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance_type</td>
<td>Sets the type of the convergence tolerance. Can be absolute or relative. Default Behavior “relative”</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance_target</td>
<td>Sets the target of the MLMC sample allocation, i.e. the constraint of the MLMC sample allocation problem Default Behavior “variance_-constraint”</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Stopping criterion based on number of refinement iterations within the multilevel sample allocation</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td>Stopping criterion based on maximum function evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
</tbody>
</table>
Description

An adaptive sampling method that utilizes multilevel relationships within a hierarchical surrogate model in order to improve efficiency through variance reduction.

In the case of a multilevel relationship, multilevel Monte Carlo methods are used to compute an optimal sample allocation per level.

Multilevel Monte Carlo

The Monte Carlo estimator for the mean is defined as

$$E[Q] ≡ \hat{Q}^{MC} = \frac{1}{N} \sum_{i=1}^{N} Q^{(i)}$$

In a multilevel method with $L$ levels, we replace this estimator with a telescoping sum:

$$E[Q] ≡ \hat{Q}^{ML} = \sum_{l=0}^{L} \frac{1}{N_l} \sum_{i=1}^{N_l} (Q^{(i)}_l - Q^{(i)}_{l-1}) \equiv \sum_{l=0}^{L} \hat{Y}_l^{MC}$$

This decomposition forms discrepancies for each level greater than 0, seeking reduction in the variance of the discrepancy $Y$ relative to the variance of the original response $Q$. The number of samples allocated for each level ($N_l$) is based on a total cost minimization procedure that incorporates the relative cost and observed variance for each of the $Y_l$.

Default Behavior

The \texttt{multilevel_sampling} method employs Monte Carlo sample sets by default, but this default can be overridden to use Latin hypercube sample sets using \texttt{sample_type lhs}.

Expected Output

The \texttt{multilevel_sampling} method reports estimates of the first four moments and a summary of the evaluations performed for each model fidelity and discretization level. The method does not support any level mappings (response, probability, reliability, generalized reliability) at this time.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the \texttt{hdf5} keyword, this method writes the following results to HDF5:

- **Sampling Moments** (moments only, not confidence intervals)

In addition, the execution group has the attribute \texttt{equiv_hf_evals}, which records the equivalent number of high-fidelity evaluations.

Usage Tips

The multilevel sampling method must be used in combination with a hierarchical model specification, and supports either a sequence of model forms or a sequence of discretization levels. For the former, each model form must provide a scalar \texttt{solution_level_cost} and for the latter, it is necessary to identify the variable string descriptor that controls the resolution levels using \texttt{solution_level_control} as well as the associated array of relative costs using \texttt{solution_level_cost}.
Examples

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
    pilot_samples = 20  seed = 1237
    max_iterations = 10
    convergence_tolerance = .001
```

does not specify a multilevel Monte Carlo study in combination with the model identified by the HIERARCH pointer. This
model specification provides a one-dimensional hierarchy, typically defined by a single model fidelity with multiple
discretization levels, but may also be provided as multiple ordered model fidelities, each with a single (or default)
discretization level. An example of the former (single model fidelity with multiple discretization levels) follows:

```plaintext
model,
  id_model = 'HIERARCH'
  surrogate hierarchical
  ordered_model_fidelities = 'SIM1'
  correction additive zeroth_order

model,
  id_model = 'SIM1'
  simulation
    solution_level_control = 'N_x'
    solution_level_cost = 630. 1260. 2100. 4200.
```

Refer to dakota/test/dakota_uq_+mlmc.in in the source distribution for additional examples.

See Also

These keywords may also be of interest:

- multifidelity_sampling
- multilevel_multifidelity_sampling

7.2.57.1 seed_sequence

Sequence of seed values for multi-stage random sampling

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** system-generated (non-repeatable)
Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 100 50 40 30 20
    seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

See Also

These keywords may also be of interest:

- seed
- fixed_seed

7.2.57.2 fixed_seed

- Keywords Area
- method
- multilevel_sampling
- fixed_seed

Reuses the same seed value for multiple random sampling sets
7.2 method

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed

7.2.57.3  pilot_samples

• Keywords Area
• method
• multilevel_sampling
• pilot_samples
```

Initial set of samples for multilevel sampling methods.

Specification

Alias: initial_samples
Argument(s): INTEGERLIST

Description

The pilot sample provides initial estimates of variance and/or correlation within the first iteration of a multilevel and/or control variate approach.

Default Behavior

100 samples per model fidelity and/or discretization level.

Usage Tips

The number of specified values can be none (default values used for all fidelities and levels), one (all fidelities and levels use the same specified value), the number of discretization levels for every model (each model uses the
same discretization level profile), or the aggregate number of discretization levels for all models (samples for each discretization level are distinct for each model).

7.2.57.4 sample_type

- Keywords Area

- method

- multilevel_sampling

- sample_type

Selection of sampling strategy

Specification

**Alias:** none

**Argument(s):** none

**Default:** random

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```
7.2.57.4.1 lhs

- Keywords Area
- method
- multilevel_sampling
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

**Alias:** none

**Argument(s):** none

Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.57.4.2 random

- Keywords Area
- method
- multilevel_sampling
- sample_type
- random

Uses purely random Monte Carlo sampling to sample variables
Specification

**Alias:** none  
**Argument(s):** none

**Description**

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: `multilevel_sampling`). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```plaintext
method
  sampling
    sample_type random
    samples = 200
```

7.2.57.5  `export_sample_sequence`

- **Keywords Area**
- **method**
- **multilevel_sampling**
- **export_sample_sequence**

Enable export of multilevel/multifidelity sample sequences to individual files

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional</strong> <em>(Choose One)</em></td>
<td>Tabular Format <em>(Group 1)</em></td>
<td><code>custom_annotated</code></td>
<td>Export sample sequences enabling file format customization</td>
</tr>
<tr>
<td>annotated</td>
<td>Export sample sequences with descriptors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-----------</td>
<td>----------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Export sample sequences without heading descriptors</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

When this option is active, separate output files are written for each unique sample increment and are tagged by algorithm type, simulation interface, iteration count, level count, and the number of samples as described below. The data content is comprised of the input variables only, without corresponding responses, as an intended use case is to support evaluation of these sample sets offline.

**Default Behavior**

If not specified, the **annotated** format is assumed.

**Expected Output**

Separate output files are generated according to the following format: `{ml/cv}_{interface_id}_{iteration_number}_{level_number}_{number_of_samples}.dat`.

With respect to the algorithm type, interface id, and level counter, the following definitions are employed:

- **Multilevel MC with hierarchical model**: `ml_` is pre-pended for all sample increments and sample sets are tagged with the interface id from the HF model.
- **Control Variate MC with hierarchical model (deprecated)**: `cv_` is prepended for all sample increments, shared sample sets are tagged with the HF interface id, and LF-only refinements are tagged by the LF interface id.
- **Multilevel-Control Variate MC with hierarchical model**: employs a combination of exports from the two cases above. `ml_` is pre-pended for all HF sample increments and sample sets are tagged with the interface id from the HF model; `cv_` is prepended for all LF increments and sample sets are tagged with the LF interface id. Note that the LF model shares the same `ml_` sample sets, but a redundant file is not created for this data.
- **Non-hierarchical approaches (Multifidelity MC, Approximate Control Variate)**: `cv_` is prepended for all sample increments and sample sets are tagged with the corresponding model interface id. The level number corresponds to the index of the approximation model for LF increments and to the id of the truth model (number of LF approximations + 1) for shared increments.

With respect to iteration count, pilot samples are tagged with iteration 0.

**Examples**

The following method block

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  output silent
  export_sample_sequence

```

results in enabling the sample output of sample increments for each level to individual files using the default **annotated** format.

The following variables block
variables,
  id_variables = 'LF_VARS'
uniform_uncertain = 7
  lower_bounds = 7 * -1.
  upper_bounds = 7 * 1.
  descriptors = 'u1' 'u2' 'u3' 'u4' 'u5' 'u6' 'u7'
descriptors = 2
  num_set_values = 4
  set_values = 5 15 30 60 # number of spatial coords
  initial_state = 5 3 # number of Fourier solution modes
  descriptors = 'N_x' 'N_mod'

illustrates how to define descriptors for the variables. For this case, the descriptors u1 through u7, N_x, and N_mod are reported in the sample files to help annotate the data.

7.2.57.5.1 custom_annotated

  • Keywords Area
  • method
  • multilevel_sampling
  • export_sample_sequence
  • custom_annotated

Export sample sequences enabling file format customization

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
7.2 method

Description

Sample sequences are written on file with a user-defined format. If the keyword header is present the heading is used, if the keyword eval_id is present the sample number (per level) is included and if interface_id is provided then the interface identification is reported.

Examples

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  output silent
  export_sample_sequence custom_annotated eval_id
```

results in enabling the sample output on file with a customized format including only the progressive number of the sample (eval_id).

7.2.57.5.1.1 header

- Keywords Area
- method
- multilevel_sampling
- export_sample_sequence
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.57.5.1.2 eval_id

- Keywords Area
- method
- multilevel_sampling
- export_sample_sequence
- custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

*See description of parent* custom_annotated

7.2.57.1.3  interface_id

- Keywords Area
- method
- multilevel_sampling
- export_sample_sequence
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

*See description of parent* custom_annotated

7.2.57.2  annotated

- Keywords Area
- method
- multilevel_sampling
- export_sample_sequence
- annotated

Export sample sequences with descriptors

**Specification**

**Alias:** none  
**Argument(s):** none

**Default:** annotated
7.2 method

Description

Sample sequences are written on file using header descriptors for the data set and a per-level progressive number, eval_id. The interface identifier, interface_id, is also reported as well as the model descriptors.

7.2.5  freeform

- Keywords Area
- method
- multilevel_sampling
- export_sample_sequence
- freeform

Export sample sequences without heading descriptors

Specification

Alias: none
Argument(s): none
Default: annotated

Description

Sample sequences are written on file without using any heading descriptors for the data set. The interface identifier, interface_id, and the model descriptors are still reported.

7.2.6  allocation_target

- Keywords Area
- method
- multilevel_sampling
- allocation_target

Allocation statistics/target for the MLMC sample allocation.

Specification

Alias: none
Argument(s): none
Default: mean

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Allocation target (Group 1)</td>
<td>mean</td>
<td>Fit MLMC sample allocation to control the variance of the estimator for the mean.</td>
</tr>
</tbody>
</table>

### Description

Computes the variance of the estimator for a given statistics/target and fits sample allocation by solving the corresponding optimization problem. The `allocation_target` specification supports options of `mean` (default), `variance` an analytic approximation is available. Additionally, a numerical optimization can be used in that case, see `variance` option optimization.

**Default Behavior** "mean"

### Examples

The following method block

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = variance
```

uses the variance as sample allocation target by computing its variance.

7.2.57.6.1 mean

- Keywords Area
- method
- multilevel_sampling
- allocation_target
- mean

Fit MLMC sample allocation to control the variance of the estimator for the mean.
7.2 method

Specification

Alias: none

Argument(s): none

Description

Computes the variance of the mean estimator and fits sample allocation by solving the corresponding optimization problem. This optimization problem is solved in analytically closed form.

Examples

The following method block

```plaintext
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 20 seed = 1237
    convergence_tolerance = .01
    allocation_target = mean
```

uses the mean as sample allocation target by computing its variance.

7.2.57.6.2 variance

- Keywords Area
- method
- multilevel_sampling
- allocation_target
- variance

Fit MLMC sample allocation to control the variance of the estimator for the variance.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>optimize</td>
<td>Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the variance.</td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

Computes the variance of the sampling estimator for the variance and fits sample allocation by solving the corresponding optimization problem. This optimization problem is obtained in closed form with an analytical approximation. Additionally, a numerical optimization can be used in that case, see optimization.

Examples

The following method block

```dakota
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 20 seed = 1237
    convergence_tolerance = .01
    allocation_target = variance
```

uses the variance as sample allocation target by computing its variance.

Theory

A single level unbiased estimator for the variance of a generic QoI at the highest level $M_L$ of the hierarchy can be written as

$$\text{Var}[Q_{M_L}] \approx \frac{1}{N_{M_L}-1} \sum_{i=1}^{N_{M_L}} \left( Q_{M_L}^{(i)} - \mathbb{E}[Q_{M_L}] \right)^2.$$

The multilevel extension for this estimator is obtained by writing

$$\text{Var}[Q_{L}] \approx \hat{Q}_{L,2}^L = \sum_{\ell=0}^{L} \hat{Q}_{\ell,2} - \hat{Q}_{\ell-1,2},$$

where

$$\hat{Q}_{\ell,2} = \frac{1}{N_{\ell} - 1} \sum_{i=1}^{N_{\ell}} \left( Q_{\ell}^{(i)} - \hat{Q}_{\ell} \right)^2 \quad \text{and} \quad \hat{Q}_{\ell-1,2} = \frac{1}{N_{\ell-1} - 1} \sum_{i=1}^{N_{\ell-1}} \left( Q_{\ell-1}^{(i)} - \hat{Q}_{\ell-1} \right)^2.$$

As for the expected value case, we want to obtain an optimal sample allocation per level that minimizes the cost to obtain an estimator with a prescribed variance. The variance of the multilevel estimator $\hat{Q}_{L,2}^M$ can be written as

$$\text{Var}[\hat{Q}_{L,2}^M] \approx \frac{1}{N_{\ell} - 1} \left( \hat{Q}_{\ell,4} - \frac{N_{\ell} - 3}{N_{\ell} - 1} \left( \hat{Q}_{\ell,2} \right)^2 \right),$$

where $\hat{Q}_{\ell,4}$ denotes the sampling estimator for the fourth order central moment. For more details about the expression that each single term takes in the previous expression, please refer to the Theory Manual.

The final sample allocation is obtained by solving a minimization problem

$$\min_{N_{\ell}} \sum_{\ell=0}^{L} C_{\ell} N_{\ell} \quad \text{s.t.} \quad \text{Var}[\hat{Q}_{L,2}^M] = \varepsilon^2 / 2.$$

This optimization problem can be solved in two different ways, namely an analytical approximation and by solving a non-linear optimization problem. The analytical approximation follows the approach described in [Pisaroni2017] and introduces a helper variable

$$\hat{V}_{2,\ell} := \text{Var}[\hat{Q}_{\ell,2}] \cdot N_{\ell},$$

and the minimization problem is formulated as

$$f(N_{\ell}, \lambda) = \sum_{\ell=0}^{L} N_{\ell} \hat{V}_{\ell} + \lambda \left( \sum_{\ell=0}^{L} N_{\ell}^{-1} \hat{V}_{2,\ell} - \varepsilon^2 / 2 \right).$$
This formulation has a closed form solution (similarly to the expected value case)

\[ N_\ell = \frac{2}{\varepsilon^2} \left[ \sum_{k=0}^{L} (\hat{V}_2,\ell k_k)^{1/2} \right] \sqrt{\frac{\hat{V}_2,\ell}{\epsilon^2}}. \]

If the option optimization is specified the previous optimization problem is solved numerically via either OPTPP (default choice) or NPSOL (if available).

See Also

These keywords may also be of interest:

- optimization

7.2.57.6.2.1 optimization

- Keywords Area
- method
- multilevel_sampling
- allocation_target
- variance
- optimization

Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the variance.

Specification

Alias: none

Argument(s): none

Description

Employ numerical optimization to solve the optimization problem for finding the optimal sample allocation when targeting the variance. This option needs NPSOL (preferred choice) or OPTPP (alternative choice) installed.

Default Behavior "off"

Examples

The following method block

```
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 20 seed = 1237
    convergence_tolerance = .01
    allocation_target = variance
    optimization
```

uses the variance as sample allocation target and solves the resulting optimization problem using numerical optimization.
Fit MLMC sample allocation to control the variance of the estimator for the standard deviation.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>optimization</td>
<td>Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the standard_deviation.</td>
</tr>
</tbody>
</table>

**Description**

Computes the variance of the sampling estimator for the standard_deviation and fits sample allocation by solving the corresponding optimization problem.

**Examples**

The following method block

```plaintext
method,
  model_pointer = ‘HIERARCH’
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = standard_deviation
```

uses the standard_deviation as sample allocation target by computing its variance.

**See Also**

These keywords may also be of interest:

- variance
Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the standard_deviation.

Specification

**Alias:** none

**Argument(s):** none

**Description**

Employ numerical optimization to solve the optimization problem for finding the optimal sample allocation when targeting the standard deviation. This option needs NPSOL (preferred choice) or OPTPP (alternative choice) installed.

**Default Behavior** "off"

**Examples**

The following method block

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = standard_deviation
  optimization
```

uses the standard deviation as sample allocation target and solves the resulting optimization problem using numerical optimization.

7.2.57.6.4 scalarization

- **Keywords Area**
- **method**
- **multilevel_sampling**
- **allocation_target**
- **scalarization**

Fit MLMC sample allocation to a mixture of terms of means and standard deviations.
Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>scalarization_response_mapping</td>
<td>Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the scalarization.</td>
</tr>
</tbody>
</table>

Description

Fit MLMC sample allocation to control the variance of the estimator for a mixture of terms of means and standard deviations. The exact scalarized formulation is given by the keyword `scalarization_response_mapping`.

Examples

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = scalarization
  scalarization_response_mapping = 1 0 0 0
                                 0 0 1 3
```

uses the standard deviation as sample allocation target by computing its variance. In this example, we assume a problem with two responses where the first line in scalarization_response_mapping refers to the first response, the second line to the second response. In the first line we only use 1 times the mean as quantity of interest. For the second response, we use 1 time the mean plus 3 times the standard deviation of the second quantity of interested. This behavior mimics the keywords `primary_response_mapping` and `secondary_response_mapping`.

See Also

These keywords may also be of interest:

- variance
  7.2.57.6.4.1 scalarization_response_mapping

  - Keywords Area
  - method
  - multilevel_sampling
7.2 method

- allocation_target
- scalarization
- scalarization_response_mapping

TODO: Need a Blurb

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** default scalarized contribution to mlmc sample allocation (1 Mean, 3 Sigma)

**Description**

**Default Behavior**

**FAQ**

See also `primary_response_mapping` and `secondary_response_mapping`

7.2.57.6.4.2 optimization

- Keywords Area
- method
- multilevel_sampling
- allocation_target
- scalarization
- optimization

Solve the optimization problem for the sample allocation by numerical optimization in the case of sampling estimator targeting the scalarization.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Employ numerical optimization to solve the optimization problem for finding the optimal sample allocation when targeting the scalarization metric. This option needs NPSOL (preferred choice) or OPTPP (alternative choice) installed.

**Default Behavior** "off"
Examples

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = scalarization
    scalarization_response_mapping = 1 0 0 0
    0 0 1 3
  optimization
```

uses the scalarization as sample allocation target and solves the resulting optimization problem using numerical optimization.

See Also

These keywords may also be of interest:

- scalarization

7.2.57.7 qoi_aggregation

- Keywords Area
- method
- multilevel_sampling
- qoi_aggregation

Aggregation strategy for the QoIs statistics for problems with multiple responses in the MLMC algorithm

Specification

**Alias:** none

**Argument(s):** none

**Default:** sum

**Child Keywords:**

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<tr>
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<th>Description of Group Qoi Aggregation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>sum</td>
<td>Aggregate the variances over all QoIs to generate a target for each level in a MLMC algorithm.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>max</td>
<td>Compute sample allocation for each response and use maximum over responses for each level in a MLMC algorithm.</td>
</tr>
</tbody>
</table>
Description

In the multilevel method a variance of the allocation_target is computed for each of the responses and their levels \( Y_i^\ell, i=1,\ldots,R, \ell=0,\ldots,L \). Setting qoi_aggregation describes the rule on how to aggregate those variances over multiple response functions. Supported options are sum (default) and max. For sum, the variances are aggregated and a single sample allocation is computed. For max, an individual sample allocation for each response using the respective variances over levels is computed and the maximum over all responses for each level is taken (worst case scenario allocation).

**Default Behavior** "sum"

Examples

The following method block

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = mean
  qoi_aggregation = sum
```

uses the sum rule to aggregate the variance over the qois.

7.2.57.7.1 sum

- Keywords Area
- method
- multilevel_sampling
- qoi_aggregation
- sum

Aggregate the variances over all QoIs to generate a target for each level in a MLMC algorithm.

Specification

**Alias:** none

**Argument(s):** none

Description

The variances of the allocation_target of \( Y_i^\ell, i=1,\ldots,R \) are aggregated and a single sample allocation is computed.

Examples

The following method block

```dakota
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = mean
  qoi_aggregation = sum
```
uses the sum rule to aggregate the variance over the qois.

7.2.57.2 max

- Keywords Area
- method
- multilevel_sampling
- qoi_aggregation
- max

Compute sample allocation for each response and use maximum over responses for each level in a MLMC algorithm.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

An individual sample allocation for each response using the respective variances of the $allocation_target$ of $Y_i^\ell$ over all levels is computed; the maximum over all responses for each level is taken.

**Examples**

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 20 seed = 1237
  convergence_tolerance = .01
  allocation_target = mean
  qoi_aggregation = max
```

uses the max rule to aggregate the variance over the qois.

7.2.57.8 convergence_tolerance

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance

Stopping criterion based on relative error

**Specification**

**Alias**: none

**Argument(s)**: REAL

**Default**: 1.e-4
Description

Multilevel sampling seeks an error balance between the estimator variance and the remaining bias error at the highest level, the two contributors to mean squared error (MSE). Since the remaining bias error is generally unknown, the convergence_tolerance is used to provide a error target relative to the Multifidelity Monte Carlo estimator variance resulting from the pilot sample. If the pilot samples are not shaped for the low-fidelity model, i.e. the number of low-fidelity evaluations is equal to the number of high-fidelity evaluations for each level, the Multifidelity estimator falls back to a Multilevel Monte Carlo estimator which is used to assess the estimator pilot samples variance. The samples allocated at each level are proportional to $\frac{1}{\epsilon^2}$, so each order of magnitude reduction in convergence_tolerance will tend to increase the sample allocation by two orders of magnitude. Therefore, this control should be used with care to avoid allocation of huge sample sets that could overrun available memory.

Default Behavior

The default value for convergence_tolerance is currently .0001, which may be too resolved for expensive simulations or high variance QoI.

7.2.57.9 convergence_tolerance_type

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_type

Sets the type of the convergence tolerance. Can be absolute or relative.

Default Behavior "relative"

Specification

Alias: none

Argument(s): none

Default: relative

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Convergence tolerance type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>absolute</td>
<td>Absolute value for the convergence tolerance.</td>
</tr>
</tbody>
</table>

7.2.57.9.1 relative

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_type
- relative

Use a relative converge criterion.
Specification

Alias: none
Argument(s): none

Description

Computes an initial tolerance $\varepsilon_{init}^2$ using the pilot samples and applies the value of keyword convergence_tolerance $c_t$ to compute the target convergence_tolerance as $\varepsilon_{target}^2 = \varepsilon_{init}^2 \cdot c_t$. Similarly if convergence_tolerance_target is equal to cost_constraint. See description in current reference manual at method.multilevel_sampling.convergence_tolerance as reference as well.

7.2.57.9.2 absolute

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_type
- absolute

Absolute value for the convergence tolerance.

Specification

Alias: none
Argument(s): none

Description

Expects an absolute value for the convergence tolerance which will be used as target. If keyword convergence_tolerance_target is equal to variance_constraint, this value is a target variance ($\varepsilon^2$); if keyword convergence_tolerance_target is equal to cost_constraint, this value is a target cost.

7.2.57.10 convergence_tolerance_target

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_target

Sets the target of the MLMC sample allocation, i.e. the constraint of the MLMC sample allocation problem

Default Behavior "variance_constraint"
7.2 method

Specification

Alias: none
Argument(s): none
Default: variance_constraint
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Convergence tolerance target (Group 1)</td>
<td>variance_constraint</td>
<td>Minimizes the cost while targeting a certain variance given in keyword convergence_tolerance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cost_constraint</td>
<td>Minimizes the variance of the estimator while targeting a certain cost given in keyword convergence_tolerance.</td>
</tr>
</tbody>
</table>

7.2.57.10.1 variance_constraint

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_target
- variance_constraint

Minimizes the cost while targeting a certain variance given in keyword convergence_tolerance.

**Specification**

**Alias:** none  
**Argument(s):** none

7.2.57.10.2 cost_constraint

- Keywords Area
- method
- multilevel_sampling
- convergence_tolerance_target
- cost_constraint

Minimizes the variance of the estimator while targeting a certain cost given in keyword convergence_tolerance.

**Specification**

**Alias:** none  
**Argument(s):** none
7.2.57.11 max_iterations

- Keywords Area
- method
- multilevel_sampling
- max_iterations

Stopping criterion based on number of refinement iterations within the multilevel sample allocation

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

**Description**

Multilevel sampling is an iterative procedure that estimates the optimal number of samples for each level based on cost and observed variance. On each iteration, additional samples are performed and more accurate variance estimates are computed, leading to updated sample allocations. The process terminates when either no additional samples are allocated or the max_iterations control is enforced.

**Default Behavior**

The default value for max_iterations varies by method. For multilevel_sampling, the default value is 25.

7.2.57.12 max_function_evaluations

- Keywords Area
- method
- multilevel_sampling
- max_function_evaluations

Stopping criterion based on maximum function evaluations

**Topics**

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1000
Description

For multilevel and multifidelity sampling methods, this specification provides a computational budget, indicating that the mean squared error (MSE) of the estimator is to be minimized within this budget. In these methods, the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error for the highest-fidelity/most-resolved model).

The budget is provided in terms of the equivalent number of high-fidelity/fine-resolution evaluations. Default value is SZ_MAX (the maximum storable size_t), which is a special value indicating that a budget has not been specified.

7.2.57.13 final_moments

- Keywords Area
- method
- multilevel_sampling
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
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<tr>
<td>Required (Choose One)</td>
<td>none</td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).
The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central

Overriding the default to none as follows:

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```

7.2.57.13.1 none

- **Keywords Area**
- **method**
- **multilevel_sampling**
- **final_moments**
- **none**

Omit moments from the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.
Examples

```plaintext
method,
  sampling
    samples = 50, seed = 1234
    response_levels = 3.6e+11, 1.2e+05, 3.5e+05
    final_moments none

7.2.57.13.2 standard
  - Keywords Area
  - method
  - multilevel_sampling
  - final_moments
  - standard

Output standardized moments and include them within the set of final statistics.

Specification

**Alias:** none

**Argument(s):** none

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50, seed = 1234
    response_levels = 3.6e+11, 1.2e+05, 3.5e+05
    final_moments standard

7.2.57.13.3 central
  - Keywords Area
  - method
  - multilevel_sampling
  - final_moments
  - central

Output central moments and include them within the set of final statistics.
7.2 method

Specification

Alias: none
Argument(s): none

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
    sampling
        samples = 50 seed = 1234
        response_levels = 3.6e+11 1.2e+05 3.5e+05
        final_moments central
```

7.2.57.14 distribution

- Keywords Area
- method
- multilevel_sampling
- distribution

Placeholder for future capabilities

Specification

Alias: none
Argument(s): none
Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>cumulative</td>
<td>complementary</td>
<td>Placeholder for future capabilities</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

At this stage only moments are computed as output in the framework of multilevel sampling techniques. No distribution capabilities are implemented yet.
At this stage only moments are computed as output in the framework of multilevel sampling techniques. No distribution capabilities are implemented yet.

At this stage only moments are computed as output in the framework of multilevel sampling techniques. No distribution capabilities are implemented yet.

Selection of a random number generator
7.2 Method

Specification

Alias: none
Argument(s): none
Default: Mersenne twister (mt19937)

Child Keywords:

<table>
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<th>Dakota Keyword</th>
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<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
</tbody>
</table>

Description

The rng keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the rng keyword must be accompanied by either rnum2 (pseudo-random numbers) or mt19937 (random numbers generated by the Mersenne twister). Otherwise, mt19937, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

```
method sampling
   sample_type lhs
   samples = 10
   seed = 98765
   rng rnum2
```

```
7.2.57.15.1 mt19937
```

• Keywords Area
• method
• multilevel_sampling
• rng
• mt19937

Generates random numbers using the Mersenne twister

Specification

Alias: none
Argument(s): none
Description

The mt19937 keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: http://en.wikipedia.org/wiki/Mersenne_twister.

Default Behavior

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended.

Examples

```
method                   sampling
     sample_type lhs
     samples = 10
     seed = 98765
     rng mt19937
```

7.2.57.15.2  rnum2

- Keywords Area
- method
- multilevel_sampling
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

Alias: none
Argument(s): none

Description

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

Examples

```
method                   sampling
     sample_type lhs
     samples = 10
     seed = 98765
     rng rnum2
```
7.2 method

7.2.57.16  model_pointer

• Keywords Area
• method
• multilevel_sampling
• model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = ‘UQ’

method
  id_method = ‘UQ’
  model_pointer = ‘SURR’
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
    sample_type lhs
distribution cumulative

model
  id_model = ‘SURR’
surrogate global,
dace_method_pointer = 'DACE'

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ‘x1’ ‘x2’

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.58 multifidelity_sampling

• Keywords Area
  • method
  • multifidelity_sampling

Multifidelity sampling methods for UQ

Specification

Alias: multifidelity_mc mfmc

Argument(s): none

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td></td>
<td>Sequence of seed values for multi-stage random sampling</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td></td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>pilot_samples</td>
<td>Initial set of samples for multi-level/multifidelity sampling methods.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>solution_mode</td>
<td>Solution mode for multi-level/multifidelity methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_sample_sequence</td>
<td>Enable export of multi-level/multifidelity sample sequences to individual files</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_tolerance</td>
<td>Stopping criterion based on relative error reduction</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_evaluations</td>
<td>Stopping criterion based on maximum function evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>final_moments</td>
<td>Output moments of the specified type and include them within the set of final statistics.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>
Description

An adaptive sampling method that utilizes multifidelity relationships in order to improve efficiency through variance reduction.

Two variants are currently supported, with the former now deprecated and to be replaced by the latter:

- In the case of a hierarchical surrogate model, the two-model approach of Ng and Willcox (2014) is supported and the two most extreme model fidelities or resolutions are employed as the truth model and approximation model.

- In the case of a non-hierarchical surrogate model, the multi-model approach of Peherstorfer et al. (2016) is supported for which all model instances can be integrated into the scheme. Both methods can be used with either a model form sequence or a resolution level sequence.

Control Variate Monte Carlo

In the case of two model fidelities (low fidelity denoted as LF and high fidelity denoted as HF), we employ a control variate approach as described in Ng and Willcox (2014):

\[
\hat{Q}_{HF}^{CV} = \hat{Q}_{HF}^{MC} - \beta (\hat{Q}_{LF}^{MC} - E[Q_{LF}])
\]

As opposed to the traditional control variate approach, we do not know \(E[Q_{LF}]\) precisely, but rather we estimate it more accurately than \(\hat{Q}_{LF}^{MC}\) based on a sampling increment applied to the LF model. This sampling increment is based again on a total cost minimization procedure that incorporates the relative LF and HF costs and the observed Pearson correlation coefficient \(\rho_{LFH}\) between \(Q_{LF}\) and \(Q_{HF}\). The coefficient \(\beta\) is then determined from the observed LF-HF covariance and LF variance.

Multifidelity Monte Carlo

This approach can be extended to a sequence of low-fidelity approximations using a recursive sampling approach as in Peherstorfer et al. (2016).

\[
\hat{Q}_{HF}^{CV} = \hat{Q}_{HF}^{MC} - \sum_{i=1}^{M} \beta_i (\hat{Q}_{LFi}^{MC} - E[Q_{LFi}])
\]

In this case, the variance in the estimate of the \(i^{th}\) control mean is reduced by the \((i + 1)^{th}\) control variate, such that the variance reduction is limited by the case of an exact estimate of the first control mean (referred to as OCV-1 in Gorodetsky et al., 2020).

Default Behavior

The multifidelity_sampling method employs Monte Carlo sample sets by default, but this default can be overridden to use Latin hypercube sample sets using sample_type lhs.

Expected Output

The multifidelity_sampling method reports estimates of the first four moments and a summary of the evaluations performed for each model fidelity and discretization level. The method does not support any level mappings (response, probability, reliability, generalized reliability) at this time.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- **Sampling Moments** (moments only, not confidence intervals)

In addition, the execution group has the attribute equiv_hf_evals, which records the equivalent number of high-fidelity evaluations.

Usage Tips
The *multifidelity_sampling* method can be used in combination with either a hierarchical or non-hierarchical model specification for either a model form sequence or a discretization level sequence. For a model form sequence, each model must provide a scalar `solution_level_cost`. For a discretization level sequence, it is necessary to identify the variable string descriptor that controls the resolution levels using `solution_level_control` as well as the associated array of relative costs using `solution_level_cost`.

The hierarchical two-model approach is a special case of the non-hierarchical multi-model approach. The latter gives identical results to the former when restricted to one approximation model; as such, the hierarchical two-model approach is deprecated.

**Examples**

We provide an example of a multifidelity Monte Carlo study using a non-hierarchical model specification employing multiple approximations.

The following method block:

```plaintext
method,
    model_pointer = 'NONHIER'
    multifidelity_sampling
    pilot_samples = 20  seed = 1237
    max_iterations = 10
    convergence_tolerance = .001
```

specifies MFMC in combination with the model identified by the NONHIER pointer.

This NONHIER model specification provides a one-dimensional sequence, here defined by a single truth model and a set of unordered approximation models, each with a single (or default) discretization level:

```plaintext
model,
    id_model = 'NONHIER'
    surrogate non hierarchical
    truth_model = 'HF'
    unordered_model_fidelities = 'LF1' 'LF2'

model,
    id_model = 'LF1'
    interface_pointer = 'LF1_INT'
    simulation
    solution_level_cost = 0.01

model,
    id_model = 'LF2'
    interface_pointer = 'LF2_INT'
    simulation
    solution_level_cost = 0.1

model,
    id_model = 'HF'
    interface_pointer = 'HF_INT'
    simulation
    solution_level_cost = 1.
```

Refer to `dakota/test/dakota_uq_*_cvmc.in` and `dakota/test/dakota_uq_*_mfmc.in` in the source distribution for additional examples.

**See Also**

These keywords may also be of interest:

- `multilevel_sampling`
- `multilevel_multifidelity_sampling`
- `approximate_control_variate`
7.2.58.1 seed_sequence

- Keywords Area
- method
- multifidelity_sampling
- seed_sequence

Sequence of seed values for multi-stage random sampling

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** system-generated (non-repeatable)

Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips

For $L$ levels in a multilevel method, the `seed_sequence` can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When `fixed_seed` is unspecified, the `seed_sequence` specification does not extend to the refinement of the sample set. For example, the value from the `seed_sequence` defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 100 50 40 30 20
  seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.
See Also

These keywords may also be of interest:

- `seed`
- `fixed_seed`

### 7.2.58.2 `fixed_seed`

- Keywords Area
- `method`
- `multifidelity_sampling`
- `fixed_seed`

Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run

**Description**

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

### 7.2.58.3 `pilot_samples`

- Keywords Area
- `method`
- `multifidelity_sampling`
- `pilot_samples`

Initial set of samples for multilevel/multifidelity sampling methods.
Specification

**Alias:** initial_samples

**Argument(s):** INTEGERLIST

Description

The pilot sample provides initial estimates of variance and/or correlation within the first iteration of a multi-level/multifidelity sampling approach. These initial estimates then guide the algorithm toward additional sample increments in order to iteratively achieve a prescribed error at minimum total cost, or alternatively, minimum error for a prescribed budget.

Default Behavior

100 samples per model fidelity and/or discretization level.

Usage Tips

The number of specified values can be none (default values used for all fidelities and levels), one (all fidelities and levels use the same specified value), the number of discretization levels for every model (each model uses the same discretization level profile), or the aggregate number of discretization levels for all models (samples for each discretization level are distinct for each model).

7.2.58.4 solution_mode

- Keywords Area
- method
- multifidelity_sampling
- solution_mode

Solution mode for multilevel/multifidelity methods

Specification

**Alias:** none

**Argument(s):** none

**Default:** online pilot

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Required (Choose One)</th>
<th>Description of Group Ensemble Sampling Solution Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Description of Group Ensemble Sampling Solution Mode (Group 1)</td>
<td>online_pilot</td>
<td>Specify a solution mode that includes the pilot cost within the sample allocation logic</td>
</tr>
</tbody>
</table>
Description

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. The pilot sample can be integrated according to one of three solution modes:

- **online_pilot**: include pilot sample cost within sample allocation (default)
- **offline_pilot**: treat pilot cost as offline and exclude from sample allocation
- **pilot_projection**: only perform the pilot sample and only estimate the expected performance based on a projection of the pilot statistics (useful for method selection)

**Default Behavior** Online pilot

**7.2.58.4.1 online_pilot**

- **Keywords Area**
- **method**
- **multifidelity_sampling**
- **solution_mode**
- **online_pilot**

Specify a solution mode that includes the pilot cost within the sample allocation logic

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, the cost of pilot sampling is treated as online cost and is included within the optimal sample allocation logic. This mode is the default.
**Additional Discussion**  It is typically advantageous to start from a smaller pilot sample and then rely on the adaptive logic to increase the sample levels to the optimal level required to achieve the desired accuracy or enforce the prescribed budget. Over-estimation of the pilot, while supporting more robust estimation of correlations and variances, can hide the optimal profile and result in a non-optimal allocation of resources.

7.2.58.4.2 offline_pilot

- Keywords Area
- method
- multifidelity_sampling
- solution_mode
- offline_pilot

Specify a solution mode that excludes the pilot cost from sample allocation logic

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, the cost of pilot sampling is treated as offline cost and is excluded from the optimal sample allocation logic. This mode is typically used for generating reference performance using an "oracle" approach, i.e., specifying a large pilot sample in order to obtain best-case correlation or variance estimates prior to generating a sample profile.

7.2.58.4.3 pilot_projection

- Keywords Area
- method
- multifidelity_sampling
- solution_mode
- pilot_projection

Specify a solution mode that estimates performance based on projecting initial correlation/variance estimates from a pilot sample

**Specification**

**Alias:** none

**Argument(s):** none
7.2 Description

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, evaluations are limited to the pilot sample and these initial estimates are used to project the potential performance under the assumption that these estimates are accurate.

This mode is useful for comparing the projected performance of different estimators, which can inform method selection.

7.2.5 Sample Type

• Keywords Area
• method
• multifidelity_sampling
• sample_type

Selection of sampling strategy

Specification

Alias: none
Argument(s): none
Default: random

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

Description

The sample_type keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use samples together with refinement_samples.

Default Behavior

If the sample_type keyword is present, it must be accompanied by lhs or random. In most contexts, lhs is the default (exception: multilevel_sampling uses Monte Carlo by default).

Examples

method
Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```
**7.2 Method**

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: `multilevel_sampling`). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```plaintext
method sampling
  sample_type random
  samples = 200
```

**7.2.58.6 export_sample_sequence**

- **Keywords Area**
- **method**
- **multifidelity_sampling**
- **export_sample_sequence**

Enable export of multilevel/multifidelity sample sequences to individual files

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td><code>custom_annotated</code></td>
<td>Selects custom-annotated tabular file format</td>
</tr>
</tbody>
</table>
Description

When this option is active, separate output files are written for each unique sample increment and are tagged by algorithm type, simulation interface, iteration count, level count, and the number of samples as described below. The data content is comprised of the input variables only, without corresponding responses, as an intended use case is to support evaluation of these sample sets offline.

Default Behavior

If not specified, the annotated format is assumed.

Expected Output

Separate output files are generated according to the following format: \( {\text{ml/cv}}_{\text{interface_id}}-{\text{iteration_number}}_{\text{level_number}}_{\text{number_of_samples}}.\text{dat} \).

With respect to the algorithm type, interface id, and level counter, the following definitions are employed:

- Multilevel MC with hierarchical model: \( \text{ml}_\) is pre-pended for all sample increments and sample sets are tagged with the interface id from the HF model.

- Control Variate MC with hierarchical model (deprecated): \( \text{cv}_\) is prepended for all sample increments, shared sample sets are tagged with the HF interface id, and LF-only refinements are tagged by the LF interface id.

- Multilevel-Control Variate MC with hierarchical model: employs a combination of exports from the two cases above. \( \text{ml}_\) is pre-pended for all HF sample increments and sample sets are tagged with the interface id from the HF model; \( \text{cv}_\) is prepended for all LF increments and sample sets are tagged with the LF interface id. Note that the LF model shares the same \( \text{ml}_\) sample sets, but a redundant file is not created for this data.

- Non-hierarchical approaches (Multifidelity MC, Approximate Control Variate): \( \text{cv}_\) is prepended for all sample increments and sample sets are tagged with the corresponding model interface id. The level number corresponds to the index of the approximation model for LF increments and to the id of the truth model (number of LF approximations + 1) for shared increments.

With respect to iteration count, pilot samples are tagged with iteration 0.

Examples

The following method block

```dakota
method,
   model_pointer = 'HIERARCH'
   multilevel_sampling
   pilot_samples = 20 seed = 1237
   convergence_tolerance = .01
   output silent
   export_sample_sequence
```
results in enabling the sample output of sample increments for each level to individual files using the default annotated format.

The following variables block

```dakota
variables,
   id_variables = 'LF_VARS'
   uniform_uncertain = 7
```

lower_bounds = 7*-1.
upper_bounds = 7* 1.
descriptors ‘u1’ ‘u2’ ‘u3’ ‘u4’ ‘u5’ ‘u6’ ‘u7’
discrete_state_set
  integer = 2
  num_set_values = 4 1
  set_values = 5 15 30 60 # number of spatial coords
  3 # number of Fourier solution modes
  initial_state = 5 3
  descriptors ‘N_x’ ‘N_mod’

illustrates how to define descriptors for the variables. For this case, the descriptors u1 through u7, N_x, and N_mod are reported in the sample files to help annotate the data.

7.2.58.6.1 custom_annotated

• Keywords Area
• method
• multifidelity_sampling
• export_sample_sequence
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```ini
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.58.6.1.1 header

- Keywords Area
- method
- multifidelity_sampling
- export_sample_sequence
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none
7.2 method

Description

See description of parent custom_annotated

7.2.58.6.1.2 eval_id

- Keywords Area
- method
- multifidelity_sampling
- export_sample_sequence
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.58.6.1.3 interface_id

- Keywords Area
- method
- multifidelity_sampling
- export_sample_sequence
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated
7.2.58.2  annotated

- Keywords Area
- method
- multifidelity_sampling
- export_sample_sequence
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated_header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment

*tabular_data*

   *tabular_data_file* = 'dakota_summary.dat'
   *annotated*

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1    NO_ID  0.9  1.1  0.0002   0.26  0.76
2    NO_ID 0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3    NO_ID 0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...```

7.2.58.6.3 freeform

- **Keywords Area**
- **method**
- **multifidelity_sampling**
- **export_sample_sequence**
- **freeform**

Selects freeform file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The *annotated* format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to *annotated* format, though *freeform* remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.58.7 convergence_tolerance

• Keywords Area

• method

• multifidelity_sampling

• convergence_tolerance

Stopping criterion based on relative error reduction

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration through satisfaction of a specified accuracy at minimum total cost.

For multilevel and multifidelity sampling methods, it is a relative convergence tolerance on the mean squared error (MSE), where the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error).

The reference estimator variance is then defined as follows:
• Multilevel approaches (multilevel_sampling and multilevel_multifidelity_sampling): tolerance relative to initial multilevel Monte Carlo (MLMC) error after pilot sample

• Control variate approaches (multifidelity_sampling and approximate_control_variate): relative to Monte Carlo (MC) error from pilot sample where these are all initial estimator variances for the algorithm, recognizing that control variate approaches can extract no benefit from low-fidelity pilot samples prior to the definition of corresponding sample increments.

Permissible values are between 0 and 1, non-inclusive. Default value is 1.e-4.

7.2.58.8 max_iterations

• Keywords Area

• method

• multifidelity_sampling

• max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt , local_reliability: 25; global_(reliability , interval_est , evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations
Stopping criterion based on maximum function evaluations

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

For multilevel and multifidelity sampling methods, this specification provides a computational budget, indicating that the mean squared error (MSE) of the estimator is to be minimized within this budget. In these methods, the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error for the highest-fidelity/most-resolved model).

The budget is provided in terms of the equivalent number of high-fidelity/fine-resolution evaluations. Default value is SZ_MAX (the maximum storable size_t), which is a special value indicating that a budget has not been specified.

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:
### Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

### Examples

The following method specification overrides the default to print `central` moments and include them in the set of final statistics.

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central

Overriding the default to `none` as follows:

```plaintext
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.
```
Omit moments from the set of final statistics.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

**Examples**

```
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

**7.2.58.10.2 standard**

- Keywords Area
- method
- multifidelity_sampling
- final_moments
- standard

Select standard rejection

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The use of a standard rejection approach in the Monte Carlo Markov Chain algorithm.
Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,  
sampling  
samples = 50 seed = 1234  
response_levels = 3.6e+11 1.2e+05 3.5e+05  
final_moments central
```

**Selection of cumulative or complementary cumulative functions**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** cumulative (CDF)  
**Child Keywords:**
### Distribution

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

#### Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

#### Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

#### Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.58.11.1 cumulative

- Keywords Area
- method
- multifidelity_sampling
- distribution
- cumulative

Computes statistics according to cumulative functions

#### Specification

**Alias:** none

**Argument(s):** none
Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

method	sampling
  sample_type lhs
  samples = 10
distribution cumulative

7.2.58.11.2 complementary
  • Keywords Area
  • method
  • multifidelity_sampling
  • distribution
  • complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.
Examples

```dakota
method
    sampling
        sample_type lhs
        samples = 10
        distribution complementary
```

7.2.58.12  rng

- **Keywords Area**
- **method**
- **multifidelity_sampling**
- **rng**

Selection of a random number generator

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister ( mt19937 )

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```dakota
method
    sampling
        sample_type lhs
        samples = 10
```
seed = 98765
rng rnum2

7.2.58.12.1 mt19937

• Keywords Area
• method
• multifidelity_sampling
• rng
• mt19937

Generates random numbers using the Mersenne twister

Specification

Alias: none
Argument(s): none

Description

The mt19937 keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: http://en.wikipedia.org/wiki/Mersenne_twister.

Default Behavior

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended.

Examples

method
sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng mt19937

7.2.58.12.2 rnum2

• Keywords Area
• method
• multifidelity_sampling
• rng
• rnum2

Generates pseudo-random numbers using the Pecos package
Keywords Area

Specification

Alias: none
Argument(s): none

Description

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2

7.2.58.13 model_pointer

  • Keywords Area
  • method
  • multifidelity_sampling
  • model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

  • block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior
7.2 method

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips
When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                0.1 0.2 0.6
                0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```

7.2.59 multilevel_multifidelity_sampling

- Keywords Area
- method
- multilevel_multifidelity_sampling
Multilevel-Multifidelity sampling methods for UQ

**Specification**

**Alias:** multilevel_multifidelity_mc mlmfmc

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td></td>
<td>Sequence of seed values for multi-stage random sampling</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td></td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>pilot_samples</td>
<td></td>
<td>Initial set of samples for multi-level/multifidelity sampling methods.</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td></td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>export_sample_sequence</td>
<td></td>
<td>Enable export of multi-level/multifidelity sample sequences to individual files</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td></td>
<td>Stopping criterion based on relative error reduction</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-evaluations</td>
<td></td>
<td>Stopping criterion based on maximum function evaluations</td>
</tr>
</tbody>
</table>
### Optional final_moments
Output moments of the specified type and include them within the set of final statistics.

### Optional distribution
Selection of cumulative or complementary cumulative functions

### Optional rng
Selection of a random number generator

### Optional model_pointer
Identifier for model block to be used by a method

#### Description
An adaptive sampling method that utilizes both multilevel and multifidelity relationships within a hierarchical surrogate model in order to improve efficiency through variance reduction.

In the case of a multilevel relationship, multilevel Monte Carlo methods are used to compute an optimal sample allocation per level, and in the case of a multifidelity relationship, control variate Monte Carlo methods are used to compute an optimal sample allocation per fidelity. These two approaches can also be combined, resulting in the multilevel-multifidelity sampling approach below.

**Multilevel Control Variate Monte Carlo**

If both multilevel and multifidelity structure are included within a hierarchical model specification, then an inner control variate can be applied across two model fidelities for each level within an outer multilevel approach.

On each level, a control variate is active for the discrepancy $Y_\ell$ based on

$$Y_\ell^* = Y_\ell + \alpha_\ell \left( \hat{Y}_\ell^{LF} - E[Y_\ell^{LF}] \right),$$

where $Y_\ell^{LF} = \gamma Q_\ell^{LF} - Q_\ell^{HF}$.

The optimal parameter $\gamma_\ell$ is computed from the correlation properties between model forms and discretization levels (see the theory manual for further details) and the optimal allocation $N_\ell$ (per level) is finally obtained from it.

#### Default Behavior
The `multilevel_multifidelity_sampling` method employs Monte Carlo sample sets be default, but this default can be overridden to use Latin hypercube sample sets using `sample_type lhs`.

#### Expected Output
The `multilevel_multifidelity_sampling` method reports estimates of the first four moments and a summary of the evaluations performed for each model fidelity and discretization level. The method does not support any level mappings (response, probability, reliability, generalized reliability) at this time.

#### Expected HDF5 Output
If Dakota was built with HDF5 support and run with the `hdf5` keyword, this method writes the following results to HDF5:

- **Sampling Moments** (moments only, not confidence intervals)

In addition, the execution group has the attribute `equiv_hf_evals`, which records the equivalent number of high-fidelity evaluations.
Usage Tips

The `multilevel_multifidelity_sampling` method must be used in combination with a hierarchical model specification. The highest and lowest fidelity model must provide multiple discretization levels, for which it is necessary to identify the variable string descriptor that controls the resolution levels using `solution_level_control` as well as the associated array of relative costs using `solution_level_cost`.

Examples

The following method block

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_multifidelity_sampling
    pilot_samples = 20 seed = 1237 #s0,#s1,#s2,#s3,#p0,#p1
    convergence_tolerance = .01 #s0,#s2,#s3,#p0,#p1
```

specifies a multilevel-multifidelity Monte Carlo study in combination with the model identified by the HIERARCH pointer. This model specification provides a two-dimensional hierarchy, comprised of two model forms each with four discretization levels:

```plaintext
model,
  id_model = 'HIERARCH'
  surrogate hierarchical
    ordered_model_fidelities = 'LF' 'HF'
model,
  id_model = 'LF'
  simulation
    solution_level_control = 'N_x'
    solution_level_cost = 375. 10125. 81000. 648000.
model,
  id_model = 'HF'
  simulation
    solution_level_control = 'N_x'
    solution_level_cost = 5.67e+5 4.536e+6 2.1e+7 1.68e+8
```

Refer to `dakota/test/dakota_uq_heat_eq_mlcvmc.in` in the source distribution for this case as well as additional examples.

See Also

These keywords may also be of interest:

- `multilevel_sampling`
- `multifidelity_sampling`

7.2.59.1 seed_sequence

- `Keywords Area`
- `method`
- `multilevel_multifidelity_sampling`
- `seed_sequence`

Sequence of seed values for multi-stage random sampling
Specification

Alias: none

Argument(s): INTEGERLIST

Default: system-generated (non-repeatable)

Description

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

Default Behavior

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

Usage Tips

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

Examples

Defining a multilevel Monte Carlo in the following way

```plaintext
method,
  model_pointer = 'HIERARCH'
  multilevel_sampling
  pilot_samples = 100 50 40 30 20
  seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

See Also

These keywords may also be of interest:

- seed
- fixed_seed
Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** not fixed; pattern varies run-to-run

**Description**

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

7.2.59.3 `pilot_samples`

**Specification**

**Alias:** initial_samples  
**Argument(s):** INTEGERLIST

Initial set of samples for multilevel/multifidelity sampling methods.
Description

The pilot sample provides initial estimates of variance and/or correlation within the first iteration of a multi-level/multifidelity sampling approach. These initial estimates then guide the algorithm toward additional sample increments in order to iteratively achieve a prescribed error at minimum total cost, or alternatively, minimum error for a prescribed budget.

Default Behavior

100 samples per model fidelity and/or discretization level.

Usage Tips

The number of specified values can be none (default values used for all fidelities and levels), one (all fidelities and levels use the same specified value), the number of discretization levels for every model (each model uses the same discretization level profile), or the aggregate number of discretization levels for all models (samples for each discretization level are distinct for each model).

7.2.59.4 sample_type

- Keywords Area
  - method
    - multilevel_multifidelity_sampling
  - sample_type

Selection of sampling strategy

Specification

Alias: none
Argument(s): none
Default: random
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
<td></td>
</tr>
<tr>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
<td></td>
</tr>
</tbody>
</table>

Description

The sample_type keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use samples together with refinement_samples.

Default Behavior
If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

7.2.59.4.1 `lhs`

- Keywords Area
- method
- multilevel_multifidelity_sampling
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
```
7.2 method

- multilevel_multifidelity_sampling
- sample_type
- random

Uses purely random Monte Carlo sampling to sample variables

Specification

Alias: none
Argument(s): none

Description

The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```dakota
method
   sampling
      sample_type random
      samples = 200
```

7.2.59.5 export_sample_sequence

- Keywords Area
- method
- multilevel_multifidelity_sampling
- export_sample_sequence

Enable export of multilevel/multifidelity sample sequences to individual files

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

When this option is active, separate output files are written for each unique sample increment and are tagged by algorithm type, simulation interface, iteration count, level count, and the number of samples as described below. The data content is comprised of the input variables only, without corresponding responses, as an intended use case is to support evaluation of these sample sets offline.

#### Default Behavior

If not specified, the annotated format is assumed.

#### Expected Output

Separate output files are generated according to the following format: 
\[(\text{ml/cv})\_{\text{interface\_id}}\_{\text{\textbackslash{}iteration\_number}}\_{\text{\textbackslash{}level\_number}}\_{\text{\textbackslash{}number\_of\_samples}}\].dat.

With respect to the algorithm type, interface id, and level counter, the following definitions are employed:

- **Multilevel MC with hierarchical model**: ml\_ is pre-pended for all sample increments and sample sets are tagged with the interface id from the HF model.

- **Control Variate MC with hierarchical model (deprecated)**: cv\_ is prepended for all sample increments, shared sample sets are tagged with the HF interface id, and LF-only refinements are tagged by the LF interface id.

- **Multilevel-Control Variate MC with hierarchical model**: employs a combination of exports from the two cases above. ml\_ is pre-pended for all HF sample increments and sample sets are tagged with the interface id from the HF model; cv\_ is prepended for all LF increments and sample sets are tagged with the LF interface id. Note that the LF model shares the same ml\_ sample sets, but a redundant file is not created for this data.

- **Non-hierarchical approaches (Multifidelity MC, Approximate Control Variate)**: cv\_ is prepended for all sample increments and sample sets are tagged with the corresponding model interface id. The level number corresponds to the index of the approximation model for LF increments and to the id of the truth model (number of LF approximations + 1) for shared increments.

With respect to iteration count, pilot samples are tagged with iteration 0.

#### Examples

The following method block

```plaintext
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 20 seed = 1237
    convergence_tolerance = .01
    output silent
    export_sample_sequence
```

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
results in enabling the sample output of sample increments for each level to individual files using the default annotated format.

The following variables block

```plaintext
variables,
   id_variables = 'LF_VARS'
   uniform_uncertain = 7
   lower_bounds  = 7*-1.
   upper_bounds  = 7* 1.
   descriptors 'u1' 'u2' 'u3' 'u4' 'u5' 'u6' 'u7'
   discrete_state_set
      integer = 2
      num_set_values = 4 1
      set_values = 5 15 30 60 # number of spatial coords
      3 # number of Fourier solution modes
      initial_state = 5 3
   descriptors 'N_x' 'N_mod'
```

illustrates how to define descriptors for the variables. For this case, the descriptors u1 through u7, N_x, and N_mod are reported in the sample files to help annotate the data.

7.2.59.5.1 custom_annotated

- Keywords Area

- method

- multilevel_multifidelity_sampling

- export_sample_sequence

- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```dakota
environment
  tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id   x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2 method

7.2.59.5.1.1 header

• Keywords Area
• method
• multilevel_multifidelity_sampling
• export_sample_sequence
• custom_annotated
• header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.59.5.1.2 eval_id

• Keywords Area
• method
• multilevel_multifidelity_sampling
• export_sample_sequence
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.59.5.1.3 interface_id

• Keywords Area
• method
• multilevel_multifidelity_sampling
Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent `custom_annotated`

7.2.59.5.2 annotated

- Keywords Area
- method
- multilevel_multifidelity_sampling
- export_sample_sequence
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include `interface_id`, specify `custom_annotated`

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading `eval_id` and `interface_id` columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

### 7.2.59.5.3 freeform

- **Keywords Area**
  - `method`
  - `multilevel_multifidelity_sampling`
  - `export_sample_sequence`
  - `freeform`

Selects freeform file format

**Topics**

This keyword is related to the topics:

- `file_formats`

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated
Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = ‘dakota_summary.dat’
  freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.59.6 convergence_tolerance

• Keywords Area

• method

• multilevel_multifidelity_sampling

• convergence_tolerance

Stopping criterion based on relative error reduction

Topics

This keyword is related to the topics:

• method_independent_controls
### 7.2 method

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.e-4

**Description**

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration through satisfaction of a specified accuracy at minimum total cost.

For multilevel and multifidelity sampling methods, it is a relative convergence tolerance on the mean squared error (MSE), where the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error).

The reference estimator variance is then defined as follows:

- **Multilevel approaches** (`multilevel_sampling` and `multilevel_multifidelity_sampling`): tolerance relative to initial multilevel Monte Carlo (MLMC) error after pilot sample

- **Control variate approaches** (`multifidelity_sampling` and `approximate_control_variate`): relative to Monte Carlo (MC) error from pilot sample where these are all initial estimator variances for the algorithm, recognizing that control variate approaches can extract no benefit from low-fidelity pilot samples prior to the definition of corresponding sample increments.

Permissible values are between 0 and 1, non-inclusive. Default value is 1.e-4.

### 7.2.59.7 max_iterations

**Keywords Area**  
**method**  
**multilevel_multifidelity_sampling**  
**max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- `method_independent_controls`

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_{reliability, interval_est, evidence} / efficient_global: 25*n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.59.8  \texttt{max\_function\_evaluations}

- Keywords Area
- method
- multilevel\_multifidelity\_sampling
- \texttt{max\_function\_evaluations}

Stopping criterion based on maximum function evaluations

Topics

This keyword is related to the topics:

- \texttt{method\_independent\_controls}

Specification

Alias: none

Argument(s): INTEGER

Default: 1000

Description

For multilevel and multifidelity sampling methods, this specification provides a computational budget, indicating that the mean squared error (MSE) of the estimator is to be minimized within this budget. In these methods, the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error for the highest-fidelity/most-resolved model).

The budget is provided in terms of the equivalent number of high-fidelity/fine-resolution evaluations. Default value is \texttt{SZ\_MAX} (the maximum storable size\_t), which is a special value indicating that a budget has not been specified.
7.2 method

7.2.59.9 final_moments

- Keywords Area
  - method
  - multilevel_multifidelity_sampling
  - final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>none</td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Select standard rejection</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The final_moments specification supports options of none, standard (default), or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

Overriding the default to none as follows:
allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
     sub_method_pointer  = 'UQ'
     primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
     secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
     sub_method_pointer  = 'UQ'
     primary_response_mapping = 1. 1. 0.
     secondary_response_mapping = 0. 0. 1.
```

7.2.59.9.1 none

• Keywords Area
• method
• multilevel_multifidelity_sampling
• final_moments
• none

Omit moments from the set of final statistics.

### Specification

**Alias:** none

**Argument(s):** none

### Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

### Examples

```plaintext
method, sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
```

7.2.59.9.2 standard

• Keywords Area
• method
7.2 method

- multilevel_multifidelity_sampling
- final_moments
- standard

Select standard rejection

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The use of a standard rejection approach in the Monte Carlo Markov Chain algorithm.

7.2.59.9.3 central

- Keywords Area  
- method  
- multilevel_multifidelity_sampling
- final_moments
- central

Output central moments and include them within the set of final statistics.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
    sampling  
        samples = 50 seed = 1234
        response_levels = 3.6e+11 1.2e+05 3.5e+05
        final_moments central
```
7.2.59.10  distribution

- Keywords Area
- method
- multilevel_multifidelity_sampling
- distribution

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

**Default Behavior**

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
  distribution cumulative
```
7.2.59.10.1 cumulative

- Keywords Area
- method
- multilevel_multifidelity_sampling
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
sampling
  sample_type lhs
  samples = 10
  distribution cumulative
```

7.2.59.10.2 complementary

- Keywords Area
- method
- multilevel_multifidelity_sampling
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none
Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

**Selection of a random number generator**

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Default**: Mersenne twister (mt19937)

**Child Keywords**:

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>RNG Algorithm (Group 1)</td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.
Default Behavior

If specified, the \texttt{rng} keyword must be accompanied by either \texttt{rnnum2} (pseudo-random numbers) or \texttt{mt19937} (random numbers generated by the Mersenne twister). Otherwise, \texttt{mt19937}, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

\begin{verbatim}
method sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
\end{verbatim}

7.2.59.11.1 mt19937

• Keywords Area
• method
• multilevel_multifidelity_sampling
• rng
• mt19937

Generates random numbers using the Mersenne twister

Specification

\textbf{Alias:} none

\textbf{Argument(s):} none

Description

The \texttt{mt19937} keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: \url{http://en.wikipedia.org/wiki/Mersenne_twister}.

Default Behavior

\texttt{mt19937} is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended.

Examples

\begin{verbatim}
method sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
\end{verbatim}
Generates pseudo-random numbers using the Pecos package

Specification

Alias: none
Argument(s): none

Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.59.12  model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer
**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```dakota
environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
        response_levels = 0.1 0.2 0.6
        0.1 0.2 0.6
        0.1 0.2 0.6
        sample_type lhs
        distribution cumulative

model
    id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
    id_method = 'DACE'
    model_pointer = 'DACE_M'
    sampling sample_type lhs
        samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
    id_interface = 'I1'
    system asynch evaluation_concurrency = 5
```
analysis_driver = 'text_book'
responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.60  approximate_control_variate

  • Keywords Area
  • method
  • approximate_control_variate

Approximate control variate (ACV) sampling methods for UQ

Specification

Alias: acv_sampling

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Required (Choose One)</th>
<th>Description of Group Solution Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
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<tr>
<td>Required</td>
<td></td>
<td>Approximate control variate (ACV) algorithm that employs independent samples per model</td>
<td>acv_independent_sampling</td>
<td>acv_independent_sampling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Approximate control variate (ACV) algorithm that mimics MFMC by employing a nested pyramid sample pattern This ACV variant uses sample set definitions that are similar to multifidelity Monte Carlo (MFMC), in that sample sets are nested with each new level adding an increment on top of the previous.</td>
<td>acv_multifidelity</td>
<td>acv_multifidelity</td>
</tr>
<tr>
<td>Optional</td>
<td>acv_kl</td>
<td>Approximate control variate (ACV) algorithm that optimizes a mix of hierarchial and non-hierarchial schemes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------</td>
<td>---------------------------------</td>
<td>----------------------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>pilot_samples</td>
<td>Initial set of samples for multi-level/multifidelity sampling methods.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>solution_mode</td>
<td>Solution mode for multi-level/multifidelity methods.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>truth_fixed_by_pilot</td>
<td>Option to suppress any increment to the number of truth samples.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Optimization Solver (Group 2)</td>
<td>sqp</td>
<td>Uses a sequential quadratic programming method for underlying optimization</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>nip</td>
<td>Uses a nonlinear interior point method for underlying optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>seed_sequence</td>
<td>Sequence of seed values for multi-stage random sampling.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td>Selection of sampling strategy</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Description**

An adaptive sampling method that utilizes multifidelity relationships in order to improve efficiency through variance reduction. It employs a non-hierarchical model to manage an unordered set of lower-fidelity approximations to a single truth model.

Compared to multifidelity Monte Carlo (MFMC), ACV relaxes the nested sampling of a recursive emulator, instead targeting the truth model's variance with each control variate pair. While the ensemble of control variates appears identical to MFMC:

\[
\hat{Q}_{HF}^{CV} = \hat{Q}_{HF}^{MC} - \sum_{i=1}^{M} \beta_i (\hat{Q}_{LF}^{MC} - \mathbb{E}[Q_{LF}])
\]

the sample patterns used for the constituent estimators differ as depicted in Gorodetsky et al. (2020), Figure 2. Two ACV variants are currently implemented, ACV-MF and ACV-IS, with ACV-KL to follow.
Default Behavior

The approximate_control_variate method employs Monte Carlo sample sets by default, but this default can be overridden to use Latin hypercube sample sets using sample_type lhs.

Expected Output

The approximate_control_variate method reports estimates of the first four moments and a summary of the evaluations performed for each model fidelity and discretization level. The method does not support any level mappings (response, probability, reliability, generalized reliability) at this time.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- **Sampling Moments** (moments only, not confidence intervals)

In addition, the execution group has the attribute equiv_hf_evals, which records the equivalent number of high-fidelity evaluations.

Usage Tips

The approximate_control_variate method must be used in combination with a non-hierarchical model specification that defines either a model form sequence or a discretization level sequence. For a model form sequence, each model must provide a scalar solution_level_cost. For a discretization level sequence, solution_level_control must identify the variable string descriptor that controls the resolution levels and the associated array of relative costs must be provided using solution_level_cost.

Examples

The following method block:

```plaintext
method,
    model_pointer = 'NONHIER'
    approximate_control_variate
        acv_mf nip
        pilot_samples = 20 seed = 1237
        max_iterations = 10
        convergence_tolerance = .001

specifies ACV-MF using the nonlinear interior point (NIP) solver in combination with the model identified by the NONHIER pointer.

This NONHIER model specification provides a one-dimensional sequence, here defined by a single truth model and a set of unordered approximation models, each with a single (or default) discretization level:

```plaintext
model,
    id_model = 'NONHIER'
    surrogate non_hierarchical
    truth_model = 'HF'
    unordered_model_fidelities = 'LF1' 'LF2'

model,
    id_model = 'LF1'
    interface_pointer = 'LF1_INT'
    simulation
        solution_level_cost = 1

model,
    id_model = 'LF2'
    interface_pointer = 'LF2_INT'
    simulation
        solution_level_cost = 16

model,
    id_model = 'HF'
    interface_pointer = 'HF_INT'
```
Refer to dakota/test/dakota_uq_diffusion_acv3_cost4.in and dakota/test/dakota_uq__tunable_acv.in in the source distribution for this case as well as additional examples. Refer to [Gorodetsky et al., JCP (408), 2020] for more detailed algorithm descriptions, theoretical considerations, and a helpful sample set diagram.

See Also

These keywords may also be of interest:

- multifidelity_sampling

### 7.2.60.1 acv_independent_sampling

- Keywords Area
- method
- approximate_control_variate
- acv_independent_sampling

Approximate control variate (ACV) algorithm that employs independent samples per model

**Specification**

**Alias:** acv_is

**Argument(s):** none

**Description**

This ACV variant uses sample set definitions that are similar to multilevel Monte Carlo (MLMC), in that independent sample sets are defined for each level and span a pair of consecutive models within a sequence.

**Theory**

Refer to[37] for a helpful sample set diagram.

### 7.2.60.2 acv_multifidelity

- Keywords Area
- method
- approximate_control_variate
- acv_multifidelity

Approximate control variate (ACV) algorithm that mimics MFMC by employing a nested pyramid sample pattern

This ACV variant uses sample set definitions that are similar to multifidelity Monte Carlo (MFMC), in that sample sets are nested with each new level adding an increment on top of the previous.
7.2 method

Specification

Alias: acv_mf
Argument(s): none

Theory

Refer to[37] for a helpful sample set diagram.

7.2.60.3 acv_kl

• Keywords Area
  • method
  • approximate_control_variate
  • acv_kl

Approximate control variate (ACV) algorithm that optimizes a mix of hierarchial and non-hierarchical schemes

Specification

Alias: acv_adaptive
Argument(s): none

Description

This ACV variant is not yet implemented. ACV-KL enumerates a set of free parameters, K and L, that mix hierarchial and non-hierarchical sampling schemes in order to obtain the best estimator for a particular budget or accuracy target.

Theory

Refer to[37] for additional details.

7.2.60.4 pilot_samples

• Keywords Area
  • method
  • approximate_control_variate
  • pilot_samples

Initial set of samples for multilevel/multifidelity sampling methods.

Specification

Alias: initial_samples
Argument(s): INTEGERLIST
Description

The pilot sample provides initial estimates of variance and/or correlation within the first iteration of a multilevel/multifidelity sampling approach. These initial estimates then guide the algorithm toward additional sample increments in order to iteratively achieve a prescribed error at minimum total cost, or alternatively, minimum error for a prescribed budget.

Default Behavior

100 samples per model fidelity and/or discretization level.

Usage Tips

The number of specified values can be none (default values used for all fidelities and levels), one (all fidelities and levels use the same specified value), the number of discretization levels for every model (each model uses the same discretization level profile), or the aggregate number of discretization levels for all models (samples for each discretization level are distinct for each model).

7.2.60.5 solution_mode

- Keywords Area
- method
- approximate_control_variate
- solution_mode

Solution mode for multilevel/multifidelity methods

Specification

Alias: none

Argument(s): none

Default: online pilot

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>Ensemble Sampling Solution Mode (Group 1)</td>
<td>online_pilot</td>
<td>Specify a solution mode that includes the pilot cost within the sample allocation logic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>offline_pilot</td>
<td>Specify a solution mode that excludes the pilot cost from sample allocation logic</td>
</tr>
</tbody>
</table>
Description

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. The pilot sample can be integrated according to one of three solution modes:

- **online_pilot**: include pilot sample cost within sample allocation (default)
- **offline_pilot**: treat pilot cost as offline and exclude from sample allocation
- **pilot_projection**: only perform the pilot sample and only estimate the expected performance based on a projection of the pilot statistics (useful for method selection)

**Default Behavior** Online pilot

**7.2.60.5.1 online_pilot**

- **Keywords Area**
- **method**
- **approximate_control_variate**
- **solution_mode**
- **online_pilot**

Specify a solution mode that includes the pilot cost within the sample allocation logic

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, the cost of pilot sampling is treated as online cost and is included within the optimal sample allocation logic. This mode is the default.

**Additional Discussion** It is typically advantageous to start from a smaller pilot sample and then rely on the adaptive logic to increase the sample levels to the optimal level required to achieve the desired accuracy or enforce the prescribed budget. Over-estimation of the pilot, while supporting more robust estimation of correlations and variances, can hide the optimal profile and result in a non-optimal allocation of resources.
Specify a solution mode that excludes the pilot cost from sample allocation logic

Specification

Alias: none

Argument(s): none

Description

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, the cost of pilot sampling is treated as offline cost and is excluded from the optimal sample allocation logic. This mode is typically used for generating reference performance using an "oracle" approach, i.e., specifying a large pilot sample in order to obtain best-case correlation or variance estimates prior to generating a sample profile.

Specify a solution mode that estimates performance based on projecting initial correlation/variance estimates from a pilot sample

Specification

Alias: none

Argument(s): none

Description

Multilevel / multifidelity sampling methods are adaptive UQ methods that utilize a pilot sample to estimate an initial set of correlations or variances, and then augment the pilot with additional sample increments to optimally allocate resources. In this mode, evaluations are limited to the pilot sample and these initial estimates are used to project the potential performance under the assumption that these estimates are accurate.

This mode is useful for comparing the projected performance of different estimators, which can inform method selection.
7.2.60.6 truth_fixed_by_pilot

- Keywords Area
- method
- approximate_control_variate
- truth_fixed_by_pilot

Option to suppress any increment to the number of truth samples

Specification

Alias: none
Argument(s): none

Description

In some cases, it may be impractical to increment an initial set of truth model samples, for example due to their extreme computational cost or due to the use of legacy data sets that can no longer be augmented. To fix the number of truth model evaluations to only those prescribed in the pilot sample, the truth_fixed_by_pilot keyword can be used.

Default Behavior

The default is to allow the number of truth samples to advance beyond those specified in the pilot, while seeking an optimal sample profile.

Additional Discussion

The optimization problem is formulated using linear constraints and a reduced set of design variables, one per approximation model.

This option is most relevant in the default online_pilot solution mode, which would normally iterate to determine the optimal number of truth samples. With truth_fixed_by_pilot, the number of truth samples is fixed within the optimization and no iteration occurs.

For the pilot_projection solution mode, the optimization formulation searches only over the approximation samples, providing an optimal projection subject to the fixed truth constraint.

The offline_pilot solution mode is not iterative and assumes that the offline pilot has accurately estimated the correlations and variances. In this case, the online profile is independent and it is not necessary to refine the truth model samples beyond the initial estimate of the optimal allocation. The truth_fixed_by_pilot specification has no additional effect beyond these existing constraints.

7.2.60.7 sqp

- Keywords Area
- method
- approximate_control_variate
- sqp

Uses a sequential quadratic programming method for underlying optimization
Specification

Alias: none
Argument(s): none
Default: sqp

Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The sqp keyword directs Dakota to use a sequential quadratic programming method to solve that problem. A sequential quadratic programming solves a sequence of linearly constrained quadratic optimization problems to arrive at the solution to the optimization problem.

7.2.60.8 nip

- Keywords Area
- method
- approximate_control_variate
- nip

Uses a nonlinear interior point method for underlying optimization

Specification

Alias: none
Argument(s): none
Default: sqp

Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The nip keyword directs Dakota to use a nonlinear interior point method to solve that problem. A nonlinear interior point method traverses the interior of the feasible region to arrive at the solution to the optimization problem.

7.2.60.9 seed_sequence

- Keywords Area
- method
- approximate_control_variate
- seed_sequence

Sequence of seed values for multi-stage random sampling
Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** system-generated (non-repeatable)

**Description**

This specification allows the stipulation of seed values (passed to a random number generator) for each of a set of stages within a multi-stage random sampling algorithm.

This is particularly useful for reusing sample sets (via restart) that were generated in some other context (e.g., from a single-stage study) within a multi-stage algorithm execution. Normally the random number sequence would continue from a seed specified for the initial sample set, making subsequent sample sets more difficult to recreate outside of their original context. With finer granularity in the seed specification for multi-stage samplers, intermediate portions of a sampling sequence can now be more readily recreated and reused.

**Default Behavior**

As for the scalar case, the default is no seed control, such that a query to a system clock will be used to randomize results.

**Usage Tips**

For $L$ levels in a multilevel method, the seed_sequence can be of any length and seeds will be assigned up to the number of values that are provided or up to the number of levels in the sequence. Thus, a seed_sequence of length 1 recovers the behavior of a scalar seed.

When fixed_seed is unspecified, the seed_sequence specification does not extend to the refinement of the sample set. For example, the value from the seed_sequence defines the initial sample set for a level, but then subsequent sample sets generated as data increments will allow the random number sequence to continue without resetting the seed value. This reflects the need to avoid reusing previous sample values when generating sample augmentations.

**Examples**

Defining a multilevel Monte Carlo in the following way

```dakota
method,
   model_pointer = 'HIERARCH'
   multilevel_sampling
   pilot_samples = 100 50 40 30 20
   seed_sequence = 1234 2345 3456 4567 5678
```

allows the reuse (via restart) of intermediate sample sets from other sources by matching the seed specifications for the targeted model resolutions.

**See Also**

These keywords may also be of interest:

- seed
- fixed_seed
Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** not fixed; pattern varies run-to-run

**Description**

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

**Default Behavior**

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

**Examples**

```
method sampling
  sample_type lhs
  samples = 10
  fixed_seed
```

### 7.2.60.11 sample_type

- [Keywords Area](#)
- [method](#)
- [approximate_control_variate](#)
- [sample_type](#)

Selection of sampling strategy

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** random  
**Child Keywords:**
## 7.2 Sample Type

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Sample Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td><strong>lhs</strong></td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>random</strong></td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

### Description

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

### Default Behavior

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

### Examples

```bash
method sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

#### 7.2.60.11.1 lhs

- Keywords Area
- method
- approximate_control_variate
- sample_type
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

### Specification

**Alias:** none

**Argument(s):** none

### Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

### Default Behavior
Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```dakota
method
sampling
  sample_type lhs
  samples = 20
```

7.2.60.11.2  random
  - **Keywords Area**
  - **method**
  - **approximate_control_variate**
  - **sample_type**
  - **random**

Uses purely random Monte Carlo sampling to sample variables

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```dakota
method
sampling
  sample_type random
  samples = 200
```
7.2.60.12 export_sample_sequence

- Keywords Area
- method
- approximate_control_variate
- export_sample_sequence

Enable export of multilevel/multifidelity sample sequences to individual files

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/(Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

When this option is active, separate output files are written for each unique sample increment and are tagged by algorithm type, simulation interface, iteration count, level count, and the number of samples as described below. The data content is comprised of the input variables only, without corresponding responses, as an intended use case is to support evaluation of these sample sets offline.

**Default Behavior**

If not specified, the annotated format is assumed.

**Expected Output**

Separate output files are generated according to the following format: `{ml/cv}_{interface_id}_{iteration_number}_{level_number}_{number_of_samples}.dat`.

With respect to the algorithm type, interface id, and level counter, the following definitions are employed:

- Multilevel MC with hierarchical model: `ml_` is pre-pended for all sample increments and sample sets are tagged with the interface id from the HF model.

- Control Variate MC with hierarchical model (deprecated): `cv_` is prepended for all sample increments, shared sample sets are tagged with the HF interface id, and LF-only refinements are tagged by the LF interface id.

- Multilevel-Control Variate MC with hierarchical model: employs a combination of exports from the two cases above. `ml_` is pre-pended for all HF sample increments and sample sets are tagged with the interface id from the HF model; `cv_` is prepended for all LF increments and sample sets are tagged with the LF interface id. Note that the LF model shares the same `ml_` sample sets, but a redundant file is not created for this data.
• Non-hierarchical approaches (Multifidelity MC, Approximate Control Variate): \texttt{cv} \_\_ is prepended for all sample increments and sample sets are tagged with the corresponding model interface id. The level number corresponds to the index of the approximation model for LF increments and to the id of the truth model (number of LF approximations + 1) for shared increments.

With respect to iteration count, pilot samples are tagged with iteration 0.

Examples

The following method block

```dakota
method,
    model_pointer = 'HIERARCH'
    multilevel_sampling
    pilot_samples = 20 seed = 1237
    convergence_tolerance = .01
    output silent
    export_sample_sequence
```

results in enabling the sample output of sample increments for each level to individual files using the default annotated format.

The following variables block

```dakota
variables,
    id_variables = 'LF_VARS'
    uniform_uncertain = 7
    lower_bounds = 7*-1.
    upper_bounds = 7* 1.
    descriptors 'u1' 'u2' 'u3' 'u4' 'u5' 'u6' 'u7'
    discrete_state_set
        integer = 2
        num_set_values = 4 1
        set_values = 5 15 30 60 # number of spatial coords
        initial_state = 5 3
        descriptors 'N_x' 'N_mod'
```

illustrates how to define descriptors for the variables. For this case, the descriptors \texttt{u1} through \texttt{u7}, \texttt{N_x}, and \texttt{N_mod} are reported in the sample files to help annotate the data.

7.2.60.12.1 custom_annotated

• Keywords Area
• method
• approximate_control_variate
• export_sample_sequence
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats
Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:
<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.60.12.1.1 header

- Keywords Area
- method
- approximate_control_variate
- export_sample_sequence
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated

7.2.60.12.1.2 eval_id

- Keywords Area
- method
- approximate_control_variate
- export_sample_sequence
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated
7.2.60.12.1.3  interface_id

- Keywords Area
- method
- approximate_control_variate
- export_sample_sequence
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

**See description of parent** custom_annotated

7.2.60.12.2  annotated

- Keywords Area
- method
- approximate_control_variate
- export_sample_sequence
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.60.12.3 freeform

- Keywords Area

- method

- approximate_control_variate

- export_sample_sequence

- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

Stopping criterion based on relative error reduction

Keywords Area

• method

• approximate_control_variate

• convergence_tolerance
Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration through satisfaction of a specified accuracy at minimum total cost.

For multilevel and multifidelity sampling methods, it is a relative convergence tolerance on the mean squared error (MSE), where the MSE is typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic error).

The reference estimator variance is then defined as follows:

- Multilevel approaches (multilevel_sampling and multilevel_multifidelity_sampling): tolerance relative to initial multilevel Monte Carlo (MLMC) error after pilot sample

- Control variate approaches (multifidelity_sampling and approximate_control_variate): relative to Monte Carlo (MC) error from pilot sample where these are all initial estimator variances for the algorithm, recognizing that control variate approaches can extract no benefit from low-fidelity pilot samples prior to the definition of corresponding sample increments.

Permissible values are between 0 and 1, non-inclusive. Default value is 1.e-4.

7.2.60.14 max_iterations

- Keywords Area

- method

- approximate_control_variate

- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.60.15 \texttt{max\_function\_evaluations}

- Keywords Area
- method
- \texttt{approximate\_control\_variate}
- \texttt{max\_function\_evaluations}

Stopping criterion based on maximum function evaluations

Topics

This keyword is related to the topics:

- \texttt{method\_independent\_controls}

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

For multilevel and multifidelity sampling methods, this specification provides a computational budget, indicating that the mean squared error (MSE) of the estimator is to be minimized within this budget. In these methods, the MSE is
typically comprised of known estimator variance (stochastic error), neglecting unknown residual bias (deterministic
error for the highest-fidelity/most-resolved model).

The budget is provided in terms of the equivalent number of high-fidelity/fine-resolution evaluations. Default value
is SZ_MAX (the maximum storable size_t), which is a special value indicating that a budget has not been specified.

7.2.60.16 final_moments

- Keywords Area
- method
- approximate_control_variate
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>none</td>
<td>Omit moments from the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>standard</td>
<td>Select standard rejection</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

Description

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control
the type of these moments. The final_moments specification supports options of none, standard (default),
or central, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness,
and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case
of a nested study. For example, final_moments none can allow for a more compact specification of primary and/or secondary response mappings.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central

Overriding the default to none as follows:

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none

allows associated nested model mappings to be simplified from:

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.

to a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.


7.2.60.16.1  none

  • Keywords Area
  • method
  • approximate_control_variate
  • final_moments
  • none

Omit moments from the set of final statistics.

Specification

Alias: none

Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments none
Select standard rejection

Specification

**Alias**: none
**Argument(s)**: none

Description

The use of a standard rejection approach in the Monte Carlo Markov Chain algorithm.

Output central moments and include them within the set of final statistics.

Specification

**Alias**: none
**Argument(s)**: none

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic U-Q). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
sampling
  samples = 50 seed = 1234
  response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```
7.2 method

7.2.60.17 distribution

• Keywords Area

• method

• approximate_control_variate

• distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

method
  sampling
    sample_type lhs
    samples = 10
distribution cumulative
7.2.60.17.1 cumulative

- Keywords Area
- method
- approximate_control_variate
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method sampling
  sample_type lhs
  samples = 10
  distribution cumulative
```

7.2.60.17.2 complementary

- Keywords Area
- method
- approximate_control_variate
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none
Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```
method
    sampling
      sample_type lhs
      samples = 10
      distribution complementary
```

7.2.6.0.18  rng

- Keywords Area
- method
- approximate_control_variate
- rng

Selection of a random number generator

Specification

Alias: none

Argument(s): none

Default: Mersenne twister ( mt19937 )

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
</tbody>
</table>

Description

The rng keyword is used to indicate a choice of random number generator.
Default Behavior

If specified, the rng keyword must be accompanied by either rnum2 (pseudo-random numbers) or mt19937 (random numbers generated by the Mersenne twister). Otherwise, mt19937, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

```
7.2.60.18.1   mt19937
  • Keywords Area
  • method
  • approximate_control_variate
  • rng
  • mt19937

Generates random numbers using the Mersenne twister

Specification

Alias: none
Argument(s): none

Description

The mt19937 keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: http://en.wikipedia.org/wiki/Mersenne_twister.

Default Behavior

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjuction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```
7.2.60.18.2 rnum2

- Keywords Area
- method
- approximate_control_variate
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

**Alias**: none

**Argument(s)**: none

**Description**

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.60.19 model_pointer

- Keywords Area
- method
- approximate_control_variate
- model_pointer

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- block_pointer
Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The \texttt{model\_pointer} is used to specify which \texttt{model} block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a \texttt{model} block in the Dakota input file that has a corresponding \texttt{id\_model} with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a \texttt{model\_pointer} for each method is imperative.

See \texttt{block\_pointer} for details about pointers.

Examples

```dakota
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE\_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE\_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
```
7.2 method

    analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.61 importance_sampling

• Keywords Area
• method
• importance_sampling

Importance sampling

Topics

This keyword is related to the topics:

• uncertainty_quantification
• aleatory_uncertainty_quantification_methods
• sampling

Specification

Alias: nond_importance_sampling

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>samples</td>
<td>Number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Importance Sampling Approach (Group 1)</td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>
The **importance_sampling** method is based on ideas in reliability modeling.

An initial Latin Hypercube sampling is performed to generate an initial set of samples. These initial samples are augmented with samples from an importance density as follows:

<table>
<thead>
<tr>
<th><strong>mm_adapt_import</strong></th>
<th>Importance sampling option for probability refinement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>refinement_-samples</td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_-levels</td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
</tr>
</tbody>
</table>

- **refinement_-samples**: Number of samples used to refine a probability estimate or sampling design.
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods.
- **convergence_-tolerance**: Stopping criterion based on objective function or statistics convergence.
- **response_levels**: Values at which to estimate desired statistics for each response.
- **probability_levels**: Specify probability levels at which to estimate the corresponding response value.
- **gen_reliability_-levels**: Specify generalized reliability levels at which to estimate the corresponding response value.
- **distribution**: Selection of cumulative or complementary cumulative functions.
- **rng**: Selection of a random number generator.
- **model_pointer**: Identifier for model block to be used by a method.
• The variables are transformed to standard normal space.
• In the transformed space, the importance density is a set of normal densities centered around points which are in the failure region.
• Note that this is similar in spirit to the reliability methods, in which importance sampling is centered around a Most Probable Point (MPP).
• In the case of the LHS samples, the importance sampling density will simply be a mixture of normal distributions centered around points in the failure region.

**Options**

Choose one of the importance sampling options:

- import
- adapt_import
- mm_adapt_import

The options for importance sampling are as follows: import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). adapt_import is the same as import but is performed iteratively until the failure probability estimate converges. mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for "representative" points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).

**Theory**

Importance sampling is a method that allows one to estimate statistical quantities such as failure probabilities (e.g. the probability that a response quantity will exceed a threshold or fall below a threshold value) in a way that is more efficient than Monte Carlo sampling. The core idea in importance sampling is that one generates samples that preferentially samples important regions in the space (e.g. in or near the failure region or user-defined region of interest), and then appropriately weights the samples to obtain an unbiased estimate of the failure probability [79].

In importance sampling, the samples are generated from a density which is called the importance density: it is not the original probability density of the input distributions. The importance density should be centered near the failure region of interest. For black-box simulations such as those commonly interfaced with Dakota, it is difficult to specify the importance density a priori: the user often does not know where the failure region lies, especially in a high-dimensional space.[81]. We have developed two importance sampling approaches which do not rely on the user explicitly specifying an importance density.

**See Also**

These keywords may also be of interest:

- adaptive_sampling
- gpais
- local_reliability
- global_reliability
- sampling
- polynomial_chaos
- stoch_collocation
7.2.61.1 samples

- Keywords Area
- method
- importance_sampling
- samples

Number of samples for sampling-based methods

Specification

Alias: initial_samples
Argument(s): INTEGER
Default: 0

Description

The samples keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least \( \text{dim}+1 \) samples should be used, where \( \text{dim} \) is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \( (\text{dim}+1)(\text{dim}+2)/2 \) samples are needed. For uncertainty quantification, we recommend at least \( 10+\text{dim} \) samples. For variance_based_decomp, we recommend hundreds to thousands of samples. Note that for variance_based_decomp, the number of simulations performed will be \( N\times(\text{dim}+2) \).

Examples

```
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.61.2 seed

- Keywords Area
- method
- importance_sampling
- seed

Seed of the random number generator
Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.61.3 import

- Keywords Area
- method
- importance_sampling
- import

Importance sampling option for probability refinement

Specification

Alias: none

Argument(s): none

Description

import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).
7.2.61.4 adapt_import

- Keywords Area
- method
- importance_sampling
- adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

7.2.61.5 mm_adapt_import

- Keywords Area
- method
- importance_sampling
- mm_adapt_import

Importance sampling option for probability refinement

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).
7.2.61.6 refinement_samples

• Keywords Area
  • method
  • importance_sampling
  • refinement_samples

Number of samples used to refine a probability estimate or sampling design.

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using unimodal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.

7.2.61.7 max_iterations

• Keywords Area
  • method
  • importance_sampling
  • max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

  • method_independent_controls

Specification

Alias: none
Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt , local_reliability: 25; global_(reliability , interval_est , evidence) / efficient_global: 25*n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed `max_iterations` iterations. In cases where there are multiple nested iterations to be controlled, `max_iterations` generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- `max_function_evaluations`

7.2.61.8 convergence_tolerance

- Keywords Area
- method
- importance_sampling
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- `method_independent_controls`

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a **relative convergence tolerance** for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.
Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- **DOT**: relative tolerance that must be satisfied for two consecutive iterations
- **NL2SOL**: See nl2sol
- **NLPQLP**: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- **NPSOL**: used as a line search tolerance, not as a relative convergence tolerance

### 7.2.61.9 response_levels

- **Keywords** Area
- method
- importance_sampling
- response_levels

Values at which to estimate desired statistics for each response

**Specification**

**Alias**: none

**Argument(s)**: REALLIST

**Default**: No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**
If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method,
  sampling,
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
    num_pairs = 3 4
    abscissas = 5 8 10 .1 .2 .3 .4
    counts = 17 21 0 12 24 12 0
    descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
```
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844608686e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.61.9.1 num_response_levels

- Keywords Area
- method
- importance_sampling
Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5

7.2.61.9.2 compute
  • Keywords Area
  • method
  • importance_sampling
  • response_levels
  • compute
```

Selection of statistics to compute at each response level

Specification

Alias: none

Argument(s): none

Default: probabilities

Child Keywords:
## Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

### Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

### Expected Output

The type of statistics specified by `compute` will be reported for each response level.

### Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

### Examples

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```

### 7.2.61.9.2.1 probabilities

- Keywords Area
- `method`
- `importance_sampling`
- `response_levels`
- `compute`
• probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```
method sampling
  sample_type random
  samples = 100 seed = 1
complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                 6.0e+04 6.5e+04 7.0e+04
                 3.5e+05 4.0e+05 4.5e+05
compute probabilities
```
Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

7.2.61.9.2.3 system

- Keywords Area
- method
- importance_sampling
- response_levels
- compute
- system

Compute system reliability (series or parallel)

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
<td></td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.61.9.2.4 series

- Keywords Area
- method
- importance_sampling
- response_levels
- compute
- system
- series

 Aggregate response statistics assuming a series system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.61.9.2.5 parallel

- Keywords Area
- method
- importance_sampling
- response_levels
- compute
- system
- parallel

 Aggregate response statistics assuming a parallel system
7.2 method

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.61.10 probability_levels

• Keywords Area
• method
• importance_sampling
• probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
<td></td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

### Probability Density Function (PDF) histograms for each response function:

#### PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.1454122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

#### PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916083900e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

#### PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

### Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.00000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.00000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.61.10.1 num_probability_levels

- Keywords Area
- method
- importance_sampling
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: probability_levels evenly distributed among response functions

Description

See parent page
Specify generalized reliability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** No CDF/CCDF response levels to compute  

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.61.11.1 num_gen_reliability_levels

- Keywords Area
- method
- importance_sampling
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response
7.2 method

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.61.12 distribution

- Keywords Area
- method
- importance_sampling
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs
Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.61.12.1 cumulative

- Keywords Area
- method
- importance_sampling
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

**Alias:** none

**Argument(s):** none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.61.12.2 complementary

- Keywords Area
- method
- importance_sampling
• distribution

• complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

7.2.61.13 rng

• Keywords Area

• method

• importance_sampling

• rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none

Default: Mersenne twister ( mt19937 )

Child Keywords:
### Description

The `rng` keyword is used to indicate a choice of random number generator.

### Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

### Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

### Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.61.13.1 `mt19937`

- Keywords Area
- method
- importance_sampling
- `rng`
- `mt19937`

Generates random numbers using the Mersenne twister

### Specification

**Alias:** none

**Argument(s):** none

### Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).
Default Behavior

\texttt{mt19937} is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
\end{verbatim}

7.2.6.13.2 \texttt{rnum2}

- Keywords Area
- \texttt{method}
- \texttt{importance_sampling}
- \texttt{rng}
- \texttt{rnum2}

Generates pseudo-random numbers using the Pecos package

Specification

\textbf{Alias}: none
\textbf{Argument(s)}: none

Description

The \texttt{rnum2} keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

\texttt{rnum2} is not used by default. To change this behavior, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended over \texttt{rnum2}.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
\end{verbatim}
7.2.61.14 model_pointer

- Keywords Area
- method
- importance_sampling
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
        response_levels = 0.1 0.2 0.6
            0.1 0.2 0.6
            0.1 0.2 0.6
        sample_type lhs
        distribution cumulative

model
    id_model = 'SURR'
method
    id_method = 'DACE'
    model_pointer = 'DACE_M'
    sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = ‘x1’ ‘x2’

interface
    id_interface = 'I1'
    system asynch evaluation_concurrency = 5
    analysis_driver = ‘text_book’

responses
    response_functions = 3
    no_gradients
    no_hessians

7.2.62 gpais

    • Keywords Area
    • method
    • gpais

Gaussian Process Adaptive Importance Sampling

Topics

This keyword is related to the topics:

    • uncertainty_quantification

Specification

Alias: gaussian_process_adaptive_importance_sampling

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>build_samples</td>
<td></td>
<td>Number of initial model evaluations used in build phase</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>-------------------------------</td>
<td>--------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_-emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_build-_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx-_points_file</td>
<td>Output file for surrogate model value evaluations</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_-levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

Description

gpais is recommended for problems that have a relatively small number of input variables (e.g. less than 10-20). This method, Gaussian Process Adaptive Importance Sampling, is outlined in the paper[? ].
This method starts with an initial set of LHS samples and adds samples one at a time, with the goal of adaptively improving the estimate of the ideal importance density during the process. The approach uses a mixture of component densities. An iterative process is used to construct the sequence of improving component densities. At each iteration, a Gaussian process (GP) surrogate is used to help identify areas in the space where failure is likely to occur. The GPs are not used to directly calculate the failure probability; they are only used to approximate the importance density. Thus, the Gaussian process adaptive importance sampling algorithm overcomes limitations involving using a potentially inaccurate surrogate model directly in importance sampling calculations.

See Also

These keywords may also be of interest:

- adaptive_sampling
- local_reliability
- global_reliability
- sampling
- importance_sampling
- polynomial_chaos
- stoch_collocation

7.2.62.1 build_samples

- Keywords Area
- method
- gpais
- build_samples

Number of initial model evaluations used in build phase

Specification

**Alias:** samples

**Argument(s):** INTEGER

Description

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of build_samples is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

Examples

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.
method
gpais
  build_samples = 100
  samples_on_emulator = 100
  max_iterations = 5
  response_levels = -1.065

7.2.62.2 seed

• Keywords Area
• method
• gpais
• seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.62.3 samples_on_emulator

• Keywords Area
• method
• gpais
7.2 method

- **samples_on_emulator**

Number of samples at which to evaluate an emulator (surrogate)

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 10000

**Description**

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

**Default Behavior**

The default number of samples is method-dependent.

**Examples**

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method
  polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.62.4 import_build_points_file

- **Keywords Area**
  - method
  - gpais
  - import_build_points_file

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
</tbody>
</table>
The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.62.4.1 custom_annotated

- **Keywords Area**
  - method
  - gpais
  - `import_build_points_file`
  - custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:
7.2.62.4.1.1 header

- Keywords Area
- method
- gpais
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

7.2.62.4.1.2 eval_id

- Keywords Area
- method
- gpais
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent custom_annotated
7.2.62.4.1.3  interface_id

- Keywords Area
- method
- gpais
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.62.4.2  annotated

- Keywords Area
- method
- gpais
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated` header `eval_id`
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID  0.9 1.1 0.0002 0.26 0.76
2   NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.62.4.3 freeform

- Keywords Area
- method
- gpais
- `import_build_points_file`
- `freeform`

Selects freeform file format

Topics

This keyword is related to the topics:

- `file_formats`
7.2 method

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.62.4.4 active_only

• Keywords Area
• method
• gpais
• import_build_points_file
• active_only

Import only active variables from tabular data file
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.62.5 export_approx_points_file

- Keywords Area
- method
- gpais
- export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file

Argument(s): STRING

Default: no point export to a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate's predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.62.5.1 custom_annotated

- Keywords Area
- method
- gpais
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.62.5.1.1 header

- Keywords Area
- method
- gpais
- export_approx_points_file
7.2 method

- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.62.5.1.2 eval_id

- Keywords Area
- method
- gpais
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.62.5.1.3 interface_id

- Keywords Area
- method
- gpais
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent *custom_annotated*

7.2.62.5.2 annotated

- Keywords Area
- method
- gpais
- `export_approx_points_file`
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated` header `eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```dakota
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
        annotated

Resulting tabular file:

```evaluation
    interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
    1 NO_ID 0.9 1.1 0.0002 0.26 0.76
    2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
    3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.62.5.3 freeform

• Keywords Area
• method
• gpais
• export_approx_points_file
• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior
The annotated format is the default for tabular export/input. To change this behavior, specify freeform in the relevant export/input context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.62.6 max_iterations

- Keywords Area
- method
- gpais
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)
Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max\_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max\_iterations} generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

- \texttt{max\_function\_evaluations}

7.2.62.7 response\_levels

- Keywords Area
- method
- \texttt{gpais}
- \texttt{response\_levels}

Values at which to estimate desired statistics for each response

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: REALLIST

\textbf{Default}: No CDF/CCDF probabilities/reliabilities to compute

\textbf{Child Keywords}:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

Description

The \texttt{response\_levels} specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior
If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

**Examples**

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2..1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n', 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u', 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w', 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h', 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

<table>
<thead>
<tr>
<th>PDF for response_fn_1:</th>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
<td></td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PDF for response_fn_2:</th>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
<td></td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>PDF for response_fn_3:</th>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
<td></td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.0000000000e-06</td>
<td></td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.62.7.1 num_response_levels

- Keywords Area
- method
- gpais
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

**Description**

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```plaintext
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```
Specification

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td><strong>probabilities</strong></td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td><strong>gen_reliabilities</strong></td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td><strong>system</strong></td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

**Description**

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then on of the following must be specified: probabilities, reliabilities, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                    6.0e+04 6.5e+04 7.0e+04
                    3.5e+05 4.0e+05 4.5e+05
  compute reliabilities
```
7.2.62.7.2.1 probabilities

- Keywords Area
- method
- gpais
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The **probabilities** keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If **response_levels** is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the **probabilities** keyword should be specified in conjunction with the **compute** keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```plaintext
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

7.2.62.7.2.2 gen_reliabilities

- Keywords Area
- method
- gpais
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels
Specification

Alias: none
Argument(s): none

Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
    compute gen_reliabilities
```

7.2.62.7.2.3 system

- Keywords Area
- method
- gpais
- response_levels
- compute
- system

Compute system reliability (series or parallel)

Specification

Alias: none
Argument(s): none
Child Keywords:
## Description

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

### 7.2.62.7.2.4 series

- **Keywords Area**
- **method**
- **gpais**
- **response_levels**
- **compute**
- **system**
- **series**

Aggregate response statistics assuming a series system

### Specification

**Alias:** none

**Argument(s):** none

### Description

See parent keyword `system` for description.

### 7.2.62.7.2.5 parallel

- **Keywords Area**
- **method**
- **gpais**
- **response_levels**
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.62.8 probability_levels

• Keywords Area
• method
• gpais
• probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none
Argument(s): REALLIST
Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td>specify which probability_levels correspond to which response</td>
<td></td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not
very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method,
  sampling,
  samples = 100 seed = 1
  complementary_distribution
  probability_levels = 1. .66 .33 0.
    .8 .5 0.
    1.3 .2 0.

variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
    num_pairs = 3 4
    abscissas = 5 8 10 .1 .2 .3 .4
    counts = 17 21 0 12 24 12 0
    descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'

interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses,
  response_functions = 3
    no_gradients
    no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>2.4334239039e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
7.2 method

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.6000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.419614379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511327775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.160383790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.870246555e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.62.8.1 num_probability_levels

- Keywords Area
- method
- gpais
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

See parent page

7.2.62.9 gen_reliability_levels

- Keywords Area
- method
- gpais
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.62.9.1 num_gen_reliability_levels

- Keywords Area
- method
- gpais
7.2 method

- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.62.10 distribution

- Keywords Area
- method
- gpais
- distribution

Selection of cumulative or complementary cumulative functions

Specification

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.
Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.62.10.1  cumulative

- Keywords Area
- method
- gpais
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

**Alias:** none

**Argument(s):** none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```
Computes statistics according to complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
  distribution complementary
```

**rng**

**Keywords Area**

**method**

**gpais**

**rng**

Selection of a random number generator
Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister ( mt19937 )

**Child Keywords:** none

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.62.11.1  mt19937

- Keywords Area
- method
- gpais
- rng
- mt19937

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none
Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```xml
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng mt19937
```

7.2.6.2.11.2 `rnum2`

- Keywords Area
- `method`
- `gpais`
- `rng`
- `rnum2`

Generates pseudo-random numbers using the Pecos package

Specification

Alias: none
Argument(s): none

Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```xml
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng rnum2
```
7.2.62.12  model_pointer

- Keywords Area
- method
- gpais
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
distribution cumulative

model
  id_model = 'SURR'
```
surrogate global,  
dace_method_pointer = 'DACE'  
polynomial quadratic  

method  
id_method = 'DACE'  
model_pointer = 'DACE_M'  
sampling sample_type lhs  
samples = 121 seed = 5034 rng rnum2

model  
id_model = 'DACE_M'  
single  
interface_pointer = 'I1'

variables  
uniform_uncertain = 2  
lower_bounds = 0. 0.  
upper_bounds = 1. 1.  
descriptors = 'x1' 'x2'

interface  
id_interface = 'I1'  
system asynch evaluation_concurrency = 5  
analysis_driver = 'text_book'

responses  
response_functions = 3  
no_gradients  
no_hessians

7.2.63 adaptive_sampling

• Keywords Area

• method

• adaptive_sampling

(Experimental) Adaptively refine a Gaussian process surrogate

Topics

This keyword is related to the topics:

• uncertainty_quantification

Specification

Alias: nond_adaptive_sampling

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>initial_samples</td>
<td>initial_samples</td>
<td>Initial number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_emulator</td>
<td>samples_on_emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
</tr>
<tr>
<td>Optional</td>
<td>fitness_metric</td>
<td>fitness_metric</td>
<td>(Experimental) Specify the fitness_metric used to select the next point</td>
</tr>
<tr>
<td>Optional</td>
<td>batch_selection</td>
<td>batch_selection</td>
<td>(Experimental) How to select new points</td>
</tr>
<tr>
<td>Optional</td>
<td>refinement_samples</td>
<td>refinement_samples</td>
<td>Number of samples used to refine a probability estimate or sampling design.</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build_points_file</td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>misc_options</td>
<td>misc_options</td>
<td>(Experimental) Hook for algorithm-specific adaptive sampling options</td>
</tr>
</tbody>
</table>
### Description

This is an experimental capability that is not ready for production use at this point. It was part of an investigation into computational topology-based approaches to feature identification and surrogate refinement.

The goal in performing adaptive sampling is to construct a surrogate model that can be used as an accurate predictor to some expensive simulation, thus it is to one's advantage to build a surrogate that minimizes the error over the entire domain of interest using as little data as possible from the expensive simulation. The adaptive part alludes to the fact that the surrogate will be refined by focusing samples of the expensive simulation on particular areas of interest rather than rely on random selection or standard space-filling techniques.

At a high-level, the adaptive sampling pipeline is a four-step process:

- Evaluate the expensive simulation (referred to as the true model) at initial sample point
  1. Fit a surrogate model
  2. Create a candidate set and score based on information from surrogate
  3. Select a candidate point to evaluate the true model
  4. Loop until done

In terms of the Dakota implementation, the adaptive sampling method currently uses Latin Hypercube sampling (LHS) to generate the initial points in Step 1 above. For Step 2, we use a Gaussian process model.

<table>
<thead>
<tr>
<th>Optional max_iterations</th>
<th>Number of iterations allowed for optimizers and adaptive UQ methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
</tr>
<tr>
<td>Optional rng</td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td>Optional model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>
The default behavior is to add one point at a time. At each iteration (e.g., each loop of Steps 2-4 above), a Latin Hypercube sample is generated (a new one, different from the initial sample) and the surrogate model is evaluated at this points. These are the candidate points that are then evaluated according to the fitness metric. The number of candidates used in practice should be high enough to fill most of the input domain: we recommend at least hundreds of points for a low-dimensional problem. All of the candidates (samples on the emulator) are given a score and then the highest-scoring candidate is selected to be evaluated on the true model.

The adaptive sampling method also can generate batches of points to add at a time using the `batch_selection` and `batch_size` keywords.

See Also

These keywords may also be of interest:

- `gpais`
- `local_reliability`
- `global_reliability`
- `sampling`
- `importance_sampling`
- `polynomial_chaos`
- `stoch_collocation`

7.2.63.1 initial_samples

Initial number of samples for sampling-based methods

Specification

**Alias:** samples

**Argument(s):** INTEGER

**Default:** 0

**Description**

The `initial_samples` keyword is used to define the number of initial samples (i.e., randomly chosen sets of variable values) at which to execute a model. The initial samples may later be augmented in an iterative process.

**Default Behavior**

By default, Dakota will use the minimum number of samples required by the chosen method.

**Usage Tips**

To obtain linear sensitivities or to construct a linear response surface, at least \( \text{dim}+1 \) samples should be used, where "\( \text{dim} \)" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least
(dim+1)(dim+2)/2 samples are needed. For uncertainty quantification, we recommend at least 10+dim samples. For `variance_based_decomp`, we recommend hundreds to thousands of samples. Note that for `variance_based_decomp`, the number of simulations performed will be N=(dim+2).

Examples

```
method
  sampling
    sample_type random
    initial_samples = 20
    refinement_samples = 5
```

### 7.2.63.2 `seed`

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **seed**

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```
7.2.63.3 samples_on_emulator

- Keywords Area
- method
- adaptive_sampling
- samples_on_emulator

Number of samples at which to evaluate an emulator (surrogate)

Specification

Alias: none
Argument(s): INTEGER
Default: 400

Description

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

Default Behavior

The default number of samples is method-dependent.

Examples

Perform 10000 samples on the PCE approximation of the true model:

```
method
polynomial_chaos
  quadrature_order  = 2
  samples_on_emulator = 10000
```

7.2.63.4 fitness_metric

- Keywords Area
- method
- adaptive_sampling
- fitness_metric

(Experimental) Specify the fitness_metric used to select the next point

Specification

Alias: none
Argument(s): none
Default: predicted_variance
Child Keywords:
7.2.63.4.1 predicted_variance

- Keywords Area
- method
- adaptive_sampling
- fitness_metric
- predicted_variance

Pick points with highest variance

**Specification**

*Alias*: none  
*Argument(s)*: none

**Description**

The predicted variance metric uses the predicted variance of the Gaussian process surrogate as the score of a candidate point. Thus, the adaptively chosen points will be in areas of highest uncertainty according to the Gaussian process model.

7.2.63.4.2 distance

- Keywords Area
- method
- adaptive_sampling
- fitness_metric
- distance

Space filling metric
Specification

**Alias:** none

**Argument(s):** none

Description

The distance metric calculates the Euclidean distance in domain space between the candidate and its nearest neighbor in the set of points already evaluated on the true model. Therefore, the most undersampled area of the domain will always be selected. Note that this is a space-filling metric.

7.2.63.4.3 gradient

- Keywords Area
- method
- adaptive_sampling
- fitness_metric
- gradient

Fill the range space of the surrogate

Specification

**Alias:** none

**Argument(s):** none

Description

The gradient metric calculates the score as the absolute value of the difference in range space (the outputs) of the two points. The output values used are predicted from the surrogate model. This method attempts to evenly fill the range space of the surrogate.

7.2.63.5 batch_selection

- Keywords Area
- method
- adaptive_sampling
- batch_selection

(Experimental) How to select new points

Specification

**Alias:** none

**Argument(s):** none

**Default:** naive

**Child Keywords:**
### Description

*adaptive_sampling* is an experimental capability that is not ready for production use at this time.

With batch or multi-point selection, the true model can be evaluated in parallel and thus increase throughput before refitting our surrogate model. This proposes a new challenge as the problem of choosing a single point and choosing multiple points off a surrogate are fundamentally different. Selecting the $n$ best scoring candidates is more than likely to generate a set of points clustered in one area which will not be conducive to adapting the surrogate.

We have implemented several strategies for batch selection of points. These are described in the User’s manual and are the subject of active research.

The *batch_selection* strategies include:

1. *naive:*
2. *distance_penalty*
3. *constant_liar*
4. *topology*

#### 7.2.63.5.1 naive

- *Keywords Area*
- *method*
- *adaptive_sampling*
- *batch_selection*
- *naive*

Take the highest scoring candidates
Keywords Area

Specification

Alias: none
Argument(s): none

Description

This strategy will select the $n$ highest scoring candidates regardless of their position. This tends to group an entire round of points in the same area.

7.2.63.5.2 distance_penalty
  - Keywords Area
  - method
  - adaptive_sampling
  - batch_selection
  - distance_penalty

Add a penalty to spread out the points in the batch

Specification

Alias: none
Argument(s): none

Description

In this strategy, the highest scoring candidate is selected and then all remaining candidates are re-scored with a distance penalization factor added in to the score.

7.2.63.5.3 topology
  - Keywords Area
  - method
  - adaptive_sampling
  - batch_selection
  - topology

In this selection strategy, we use information about the topology of the space from the Morse-Smale complex to identify next points to select.

Specification

Alias: none
Argument(s): none
Description

In this strategy we look at the topology of the scoring function and select the $n$ highest maxima in the topology. To determine local maxima, we construct the approximate Morse-Smale complex. This strategy does require the user to have the Morse-Smale package.

7.2.63.5.4 constant_liar

- Keywords Area
- method
- adaptive_sampling
- batch_selection
- constant_liar

Use information from the existing surrogate model to predict what the surrogate upgrade will be with new points.

Specification

Alias: none
Argument(s): none

Description

The strategy first selects the highest scoring candidate, and then refits the surrogate using a "lie" value at the point selected. The 'lie' value is based on the surrogate predictions and not the simulation. This process repeats until $n$ points have been selected whereupon the lie values are removed from the surrogate and the selected points are evaluated on the true model and the surrogate is refit with these values.

7.2.63.6 refinement_samples

- Keywords Area
- method
- adaptive_sampling
- refinement_samples

Number of samples used to refine a probability estimate or sampling design.

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using uni-modal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial_samples in a sampling design.
7.2.63.7 import_build_points_file

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform
Examples

```python
method
def polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.63.7.1 custom_annotated

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.
Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1   x2   obj_fn   nln_ineq_con_1 nln_ineq_con_2
1   0.9  1.1   0.0002  0.26     0.76
2   0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3   0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...
```

7.2.63.7.1.1 header

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated
7.2.63.7.1.2 eval_id

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.63.7.1.3 interface_id

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated
Selects annotated tabular file format

Topics
This keyword is related to the topics:

• file_formats

Specification
Alias: none
Argument(s): none
Default: annotated format

Description
An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior
By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples
Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>@eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.63.7.3 freeform

- Keywords Area
- method
- adaptive_sampling
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• In freeform, the `num_rows x num_cols` total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

    0.9  1.1  0.0002  0.26  0.76
    0.90009  1.1  0.0001996404857  0.2601620081  0.759955
    0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.63.7.4 active_only

• Keywords Area
  • method
  • adaptive_sampling
  • import_build_points_file
  • active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
### 7.2.63.8 export_approx_points_file

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **export_approx_points_file**

Output file for surrogate model value evaluations

**Specification**

**Alias:** export_points_file  
**Argument(s):** STRING  
**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### 7.2.63.8.1 custom_annotated

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **export_approx_points_file**
- **custom_annotated**

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
7.2 method

environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...

7.2.63.8.1.1 header

- Keywords Area
- method
- adaptive_sampling
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.63.8.1.2 eval_id

- Keywords Area
- method
- adaptive_sampling
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

See description of parent custom_annotated

7.2.63.8.1.3 interface_id

• Keywords Area
• method
• adaptive_sampling
• export_approx_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.63.8.2 annotated

• Keywords Area
• method
• adaptive_sampling
• export_approx_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:
```

<table>
<thead>
<tr>
<th>%eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.63.8.3 freeform

- Keywords Area
- method
- adaptive_sampling
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specifications

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The *annotated* format is the default for tabular export/import. To change this behavior, specify *freeform* in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to *annotated* format, though *freeform* remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In *freeform*, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.900009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

**7.2.63.9** misc_options

- Keywords Area
- method
- adaptive_sampling
- misc_options

(Experimental) Hook for algorithm-specific adaptive sampling options
7.2 method

Specification

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** no misc options

**Description**

The adaptive sampling algorithm is an experimental capability and not ready for production use at this time.

7.2.63.10 max_iterations

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- **method_independent_controls**

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- **max_function_evaluations**
7.2.63.11 response_levels

- Keywords Area
- method
- adaptive_sampling
- response_levels

Values at which to estimate desired statistics for each response

**Specification**

*Alias:* none

*Argument(s):* REALLIST

*Default:* No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_levels` correspond to which response.
Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num_response_levels key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```
method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
  6.0e+04 6.5e+04 7.0e+04
  3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.63.11.1 num_response_levels

- Keywords Area
- method
- adaptive_sampling
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none
Argument(s): INTEGERLIST
**Default:** response_levels evenly distributed among response functions

**Description**

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```plaintext
method sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

### Specification

- **Alias:** none
- **Argument(s):** none
- **Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
</tbody>
</table>
Description

The `compute` keyword is used to select which forward stastical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then on of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```

7.2.63.11.2.1 probabilities

- Keywords Area
- method
- adaptive_sampling
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels
7.2 method

Specification

Alias: none
Argument(s): none

Description

The **probabilities** keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If **response_levels** is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the **probabilities** keyword should be specified in conjunction with the **compute** keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

**Examples**

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                       6.0e+04 6.5e+04 7.0e+04
                       3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

7.2.63.11.2.2  **gen_reliabilities**

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **response_levels**
- **compute**
- **gen_reliabilities**

Computes generalized reliabilities associated with response levels

**Specification**

Alias: none
Argument(s): none

Description

The **gen_reliabilities** keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.
Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```dakota
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary_distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
      6.0e+04 6.5e+04 7.0e+04
      3.5e+05 4.0e+05 4.5e+05
    compute gen_reliabilities
```

7.2.63.11.2.3 system

- **Keywords Area**
- **method**
- **adaptive_sampling**
- **response_levels**
- **compute**
- **system**

Compute system reliability (series or parallel)

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

Description

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.
For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.63.11.2.4 series

• Keywords Area
• method
• adaptive_sampling
• response_levels
• compute
• system
• series

Aggregate response statistics assuming a series system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.63.11.2.5 parallel

• Keywords Area
• method
• adaptive_sampling
• response_levels
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.
7.2.63.12 probability_levels

- Keywords Area
- method
- adaptive_sampling
- probability_levels

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Examples**

The Dakota input file below specifies a sampling method with probability levels of interest.

```
method, sampling,  
samples = 100 seed = 1  
complementary_distribution  
probability_levels = 1. .66 .33 0.  
1. .8 .5 0.  
1. .3 .2 0.  
variables,  
normal_uncertain = 2  
means = 248.89, 593.33  
std_deviations = 12.4, 29.7  
descriptors = 'TF1n' 'TF2n'
```
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'

interface,
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'
responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.1454122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239035e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.00000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.00000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.00000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.00000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.00000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.00000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.00000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.00000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.00000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.63.12.1 num_probability_levels

• Keywords Area
• method
• adaptive_sampling
• probability_levels
• num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.63.13 gen_reliability_levels

• Keywords Area
• method
• adaptive_sampling
• gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value
7.2 method

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.63.13.1 num_gen_reliability_levels

- Keywords Area
- method
- adaptive_sampling
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions
Description

See parent page

7.2.63.14  distribution

- Keywords Area
- method
- adaptive_sampling
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>cumulative</td>
<td>complementary</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

method
sampling
7.2 method

```plaintext
call test
sample_type lhs
samples = 10
distribution cumulative
```

7.2.63.14.1 cumulative

- Keywords Area
- method
- adaptive_sampling
- distribution
- cumulative

Computes statistics according to cumulative functions

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```plaintext
method
sampling
  sample_type lhs
  samples = 10
  distribution cumulative
```

7.2.63.14.2 complementary

- Keywords Area
- method
- adaptive_sampling
- distribution
- complementary

Computes statistics according to complementary cumulative functions
Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```dakota
method
    sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

```
7.2.63.15  rng
```

- Keywords Area
- method
- adaptive_sampling
- rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none

Default: Mersenne twister ( mt19937 )

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
7.2 Method

| rnum2 | Generates pseudo-random numbers using the Pecos package |

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.63.15.1 mt19937

- Keywords Area
- method
- adaptive_sampling
- rng
- mt19937

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

**Default Behavior**

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended.
Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

**7.2.63.15.2 rnum2**

- Keywords Area
- method
- adaptive_sampling
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

**7.2.63.16 model_pointer**

- Keywords Area
- method
- adaptive_sampling
- model_pointer

Identifier for model block to be used by a method
7.2 method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                  0.1 0.2 0.6
                  0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ‘x1’ ‘x2’

interface
id_interface = ’I1’
system asynch evaluation_concurrency = 5
analysis_driver = ’text_book’

responses
response_functions = 3
no_gradients
no_hessians

7.2.64 pof_darts

• Keywords Area
• method
• pof_darts

Probability-of-Failure (POF) darts is a novel method for estimating the probability of failure based on random sphere-packing.

Topics

This keyword is related to the topics:

• uncertainty_quantification

Specification

Alias: nond_pof_darts

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>build_samples</td>
<td></td>
<td>Number of initial model evaluations used in build phase</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>lipschitz</td>
<td></td>
<td>Select the type of Lipschitz estimation (global or local)</td>
</tr>
<tr>
<td>Optional</td>
<td>samples_on_-emulator</td>
<td></td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
</tr>
</tbody>
</table>
7.2.64.1 build_samples

- Keywords Area
- method
Number of initial model evaluations used in build phase

**Specification**

**Alias:** samples

**Argument(s):** INTEGER

**Description**

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of `build_samples` is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

**Examples**

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```plaintext
method gpais
  build_samples = 100
  samples_on_emulator = 100
  max_iterations = 5
  response_levels = -1.065
```

**7.2.64.2 seed**

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random `seed` control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.
If not specified, the seed is randomly generated.

**Expected Output**

If *seed* is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without *seed* specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

### 7.2.64.3 lipschitz

- **Keywords Area**
- **method**
- **pof_darts**
- **lipschitz**

Select the type of Lipschitz estimation (global or local)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Local/Global Estimate (Group 1)</td>
</tr>
<tr>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>local</td>
<td>Specify local estimation of the Lipschitz constant</td>
</tr>
<tr>
<td>global</td>
<td>Specify global estimation of the Lipschitz estimate</td>
</tr>
</tbody>
</table>

**Description**

There are two types of Lipschitz estimation used in sizing the disks used in POF Darts: *global* and *local*. The global approach uses one Lipschitz estimate for the entire domain. The local approach calculates the Lipschitz estimate separately for each Voronoi region based on nearby points. The local method is more expensive but more accurate.
7.2.64.3.1 local

    • Keywords Area
    • method
    • pof_darts
    • lipschitz
    • local

Specify local estimation of the Lipschitz constant

**Specification**

**Alias**: none
**Argument(s)**: none

**Description**

The local approach to estimate the Lipschitz constant calculates the Lipschitz estimate separately for each Voronoi region based on nearby points. The local method is more expensive but results in higher accuracy compared to the global method.

7.2.64.3.2 global

    • Keywords Area
    • method
    • pof_darts
    • lipschitz
    • global

Specify global estimation of the Lipschitz estimate

**Specification**

**Alias**: none
**Argument(s)**: none

**Description**

The global approach uses one Lipschitz estimate for the entire domain. This option is currently deactivated. Please refer to the local alternative.

7.2.64.4 samples_on_emulator

    • Keywords Area
    • method
    • pof_darts
• **samples_on_emulator**

Number of samples at which to evaluate an emulator (surrogate)

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

**Default Behavior**

The default number of samples is method-dependent.

**Examples**

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method  
  polynomial_chaos  
    quadrature_order = 2  
    samples_on_emulator = 10000
```

7.2.64.5 **response_levels**

• Keywords Area  
• method  
• pof_darts  
• response_levels

Values at which to estimate desired statistics for each response

**Specification**

**Alias:** none  
**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
</tbody>
</table>
### Description

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

### Default Behavior

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

### Expected Outputs

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

### Usage Tips

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.

### Examples

For example, specifying a `response_level` of 52.3 followed with `compute probabilities` will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the `num_response_levels` key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method,  
  sampling,  
  samples = 100 seed = 1  
  complementary distribution  
  response_levels = 3.6e+11 4.0e+11 4.4e+11  
                      6.0e+04 6.5e+04 7.0e+04  
                      3.5e+05 4.0e+05 4.5e+05  
variables,  
  normal_uncertain = 2  
  means = 248.89, 593.33  
  std_deviations = 12.4, 29.7  
  descriptors = 'TF1n', 'TF2n'  
  uniform_uncertain = 2  
  lower_bounds = 199.3, 474.63
```
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'

interface,
system async evaluation_concurrency = 5
analysis_driver = 'text_book'

responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.284460868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+01</td>
<td>5.5000000000e+01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+01</td>
<td>3.8000000000e+01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+01</td>
<td>2.3000000000e+01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e+04</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e+04</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e+02</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e+02</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.64.5.1 num_response_levels

- Keywords Area
- method
- pof_darts
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5

7.2.64.5.2 compute

- Keywords Area
- method
Selection of statistics to compute at each response level

Specification

Alias: none

Argument(s): none

Default: probabilities

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td></td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliabilities</td>
<td></td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td>system</td>
<td></td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```
method sampling
```
sample_type random
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05
compute reliabilities

7.2.64.5.2.1 probabilities
  • Keywords Area
  • method
  • pof_darts
  • response_levels
  • compute
  • probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The probabilities keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the probabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
       6.0e+04 6.5e+04 7.0e+04
      3.5e+05 4.0e+05 4.5e+05
    compute probabilities

7.2.64.5.2.2 gen_reliabilities
  • Keywords Area
  • method
Computes generalized reliabilities associated with response levels

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

**Examples**

```
method sampling  
    sample_type random  
    samples = 100 seed = 1  
    complementary distribution  
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05  
    compute gen_reliabilities
```

**7.2.64.5.2.3 system**

- **Keywords Area**
- **method**
- **pof_darts**
- **response_levels**
- **compute**
- **system**

Compute system reliability (series or parallel)

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

#### 7.2.64.5.2.4 series

- **Keywords Area**
- **method**
- **pof_darts**
- **response_levels**
- **compute**
- **system**
- **series**

Aggregate response statistics assuming a series system

#### Specification

**Alias**: none

**Argument(s)**: none

#### Description

See parent keyword `system` for description.

#### 7.2.64.5.2.5 parallel

- **Keywords Area**
- **method**
- **pof_darts**
- **response_levels**

Aggregate response statistics assuming a parallel system
Aggregate response statistics assuming a parallel system

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

See parent keyword `system` for description.

### 7.2.64.6 probability_levels

- **Keywords Area**
- **method**
- **pof_darts**
- **probability_levels**

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none  
**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not
very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1. .66 .33 0.
                     1. .8 .5 0.
                     1. .3 .2 0.
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

```
Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
       Bin Lower   Bin Upper   Density Value
---------- ---------- ---------- ---------------
2.7604749078e+11 3.4221494996e+11 5.1384774972e-12
3.4221494996e+11 4.0634975300e+11 5.1454122311e-12
4.0634975300e+11 5.4196114379e+11 2.4334239039e-12
PDF for response_fn_2:
       Bin Lower   Bin Upper   Density Value
---------- ---------- ---------- ---------------
4.643114744e+04 5.6511827775e+04 1.9839945149e-05
5.6511827775e+04 6.1603813790e+04 5.8916108390e-05
6.1603813790e+04 7.8702465755e+04 2.9242071306e-05
PDF for response_fn_3:
```
7.2 method

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.810096235e+05</td>
<td>9.0600055634e-06</td>
</tr>
<tr>
<td>3.810096235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.6000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511882775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.810096235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.6.4.6.1 num_probability_levels

- Keywords Area
- method
- pof_darts
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions
Description

See parent page

7.2.64.7 gen_reliability_levels

- Keywords Area
- method
- pof_darts
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
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<td></td>
<td>num_gen_reliability_levels</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.64.7.1 num_gen_reliability_levels

- Keywords Area
- method
- pof_darts
7.2 method

- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.64.8 distribution

- Keywords Area
- method
- pof_darts
- distribution

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

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<th>Required/Optional</th>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>complementary</td>
<td>cumulative</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The *distribution* keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.
Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```
Computes statistics according to complementary cumulative functions

Specification

**Alias**: none

**Argument(s)**: none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
  distribution complementary
```

** rng

** Keywords Area

** method

** pof_darts

** distribution

** complementary

Selection of a random number generator
Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>RNG Algorithm</strong> (Group 1)</td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```bash
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2

7.2.64.9.1 mt19937
```

**Specification**

**Alias:** none

**Argument(s):** none
7.2 method

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```plaintext
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng mt19937
```

7.2.64.9.2 rnum2

- Keywords Area
- method
- pof_darts
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

Alias: none

Argument(s): none

Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```plaintext
method sampling
  sample_type lhs
  samples = 10
  seed = 98765
  rng rnum2
```
### 7.2.64.10 model_pointer

- **Keywords Area**
- **method**
- **pof_darts**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

**Description**

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
```
7.2 method

surrogate global,
dace_method_pointer = 'DACE'

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.65 rkd_darts

  - Keywords Area
  - method
  - rkd_darts

Recursive k-d (RKD) Darts: Recursive Hyperplane Sampling for Numerical Integration of High-Dimensional Functions.

Topics

This keyword is related to the topics:

  - uncertainty_quantification

Specification

Alias: nond_rkd_darts

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>build_samples</td>
<td>Number of initial model evaluations used in build phase</td>
</tr>
</tbody>
</table>
### Keywords Area

<table>
<thead>
<tr>
<th>Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>lipschitz</td>
<td>Undocumented: Recursive k-d (RKD) Darts is an experimental capability.</td>
</tr>
<tr>
<td>samples_on_emulator</td>
<td>Number of samples at which to evaluate an emulator (surrogate)</td>
</tr>
<tr>
<td>response_levels</td>
<td>Undocumented: Recursive k-d (RKD) Darts is an experimental capability.</td>
</tr>
<tr>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
</tr>
<tr>
<td>rng</td>
<td>Undocumented: Recursive k-d (RKD) Darts is an experimental capability.</td>
</tr>
<tr>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

### Description

Disclaimer: The RKD method is currently in development mode, for further experimental verification. Please contact Dakota team if you have further questions about using this method.

Recursive k-d (RKD) darts is an algorithm to evaluate the integration of a d-dimensional black box function f(x) via recursive sampling over d, using a series of hyperplanes of variable dimensionality k = {d, d-1, ... , 0}. Fundamentally, we decompose the d-dimensional integration problem into a series of nested one-dimensional problems. That is, we start at the root level (the whole domain) and start sampling down using hyperplanes of one lower dimension, all the way down to zero (points). A d-dimensional domain is subsampled using (d-1) hyperplanes, a (d-1)-dimensional sub-domain is subsampled using (d-2) hyperplanes, and so on. Every hyperplane, regardless of its dimension, is evaluated using sampled hyperplanes of one lower dimension, as shown in the set of figures above. Each hyperplane has direct information exchange with its parent hyperplane of one higher dimension, and its children of one lower dimension.
In each one-dimensional problem, we construct a piecewise approximation surrogate model, using 1-dimensional Lagrange interpolation. Information is exchanged between different levels, including integration values, as well as interpolation and evaluation errors, in order to a) find the integration value up to that level, b) estimate the associated integration error, and c) guide the placement of future samples.

### 7.2.65.1 build_samples

- **Keywords Area**
- **method**
- **rkd_darts**
- **build_samples**

Number of initial model evaluations used in build phase

**Specification**

**Alias:** samples  
**Argument(s):** INTEGER

**Description**

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of build_samples is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

**Examples**

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```plaintext
method gpais
  build_samples = 100
  samples_on_emulator = 100
  max_iterations = 5
  response_levels = -1.065
```

### 7.2.65.2 seed

- **Keywords Area**
- **method**
- **rkd_darts**
- **seed**

Seed of the random number generator
Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347

7.2.65.3 lipschitz

• Keywords Area

• method

• rkd_darts

• lipschitz

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

Specification

Alias: none
Argument(s): none
Child Keywords:
### 7.2 method

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<td></td>
<td>global</td>
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</table>

#### 7.2.65.3.1 local

- Keywords Area
- method
- rkd_darts
- lipschitz
- local

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

**Specification**

**Alias**: none

**Argument(s)**: none

#### 7.2.65.3.2 global

- Keywords Area
- method
- rkd_darts
- lipschitz
- global

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

**Specification**

**Alias**: none

**Argument(s)**: none

#### 7.2.65.4 samples_on_emulator

- Keywords Area
- method
- rkd_darts
• **samples_on_emulator**

Number of samples at which to evaluate an emulator (surrogate)

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

How many approximate function evaluations to perform on the emulator model, e.g., to compute statistics

**Default Behavior**

The default number of samples is method-dependent.

**Examples**

Perform 10000 samples on the PCE approximation of the true model:

```plaintext
method
  polynomial_chaos
    quadrature_order = 2
    samples_on_emulator = 10000
```

7.2.65.5 **response_levels**

- **Keywords Area**

- **method**

- **rkd_darts**

- **response_levels**

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**
7.2.6.5.5.1 num_response_levels

- Keywords Area
- method
- rkd_darts
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```dakota
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```
7.2.65.5.2 compute

- Keywords Area
- method
- rkd_darts
- response_levels
- compute

Selection of statistics to compute at each response level

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

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<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td>system</td>
<td></td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

**Description**

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.
Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```

7.2.65.5.2.1 probabilities

- Keywords Area
- method
- rkd_darts
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

Specification

**Alias:** none

**Argument(s):** none

Description

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
    compute probabilities
```
Computes generalized reliabilities associated with response levels

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

**Examples**

```
method sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

**7.2.65.5.2.3 system**

```bash
Compute system reliability (series or parallel)
```

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7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<tr>
<td></td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

**Description**

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.65.5.2.4 series

- Keywords Area
- method
- rkd_darts
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

See parent keyword `system` for description.
7.2.65.5.2.5 parallel

- Keywords Area
- method
- rkd_darts
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.65.6 probability_levels

- Keywords Area
- method
- rkd_darts
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>
Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method, sampling,
samples = 100 seed = 1
complementary distribution
probability_levels = 1 .66 .33 0.
1 .8 .5 0.
1 .1 .2 0.
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
uniform_uncertain = 2
lower_bounds = 109.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u' 'TF2u'
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'
interface,
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'
responses,
response_functions = 3
no_gradients
no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:
PDF for response_fn_1:
level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.6000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.65.6.1 num_probability_levels

- Keywords Area
- method
- rkd_darts
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response
7.2 method

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.65.7 gen_reliability_levels

- Keywords Area
- method
- rkd_darts
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

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<th>Description of Group</th>
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<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
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</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
7.2.65.7.1 num_gen_reliability_levels

- Keywords Area
- method
- rkd_darts
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.65.8 distribution

- Keywords Area
- method
- rkd_darts
- distribution

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** cumulative (CDF)

**Child Keywords:**

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<tbody>
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<td></td>
<td>cumulative</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>
Description

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.65.8.1 cumulative

- Keywords Area
- method
- rkd_darts
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

**Alias**: none

**Argument(s)**: none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.
Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.65.8.2 complementary

- Keywords Area
- method
- rkd_darts
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

**Alias:** none

**Argument(s):** none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

7.2.65.9 rng

- Keywords Area
- method
- rkd_darts
- rng

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.
7.2 method

Specification

Alias: none

Argument(s): none

Default: Mersenne twister (mt19937)

Child Keywords:

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<td>Undocumented: Recursive k-d (RKD) Darts is an experimental capability.</td>
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7.2.65.9.1 mt19937

• Keywords Area
  • method
  • rkd_darts
  • rng
  • mt19937

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

Specification

Alias: none

Argument(s): none

7.2.65.9.2 rnum2

• Keywords Area
  • method
  • rkd_darts
  • rng
  • rnum2

Undocumented: Recursive k-d (RKD) Darts is an experimental capability.

Specification

Alias: none

Argument(s): none
7.2.65.10  model_pointer

- Keywords Area
- method
- rkd_darts
- model_pointer

Identifier for model block to be used by a method

Topics
This keyword is related to the topics:
- block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)

Description
The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior
If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips
When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.
See block_pointer for details about pointers.

Examples

environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
          0.1 0.2 0.6
          0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
7.2 method

surrogate global,
dace_method_pointer = 'DACE'
nominal quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.66 global_evidence

• Keywords Area

• method

• global_evidence

Evidence theory with evidence measures computed with global optimization methods

Topics

This keyword is related to the topics:

• epistemic_uncertainty_quantification_methods

• evidence_theory

Specification

Alias: nond_global_evidence

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>samples</td>
<td>Number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Solution Approach (Group 1)</td>
<td>sbo</td>
<td>Use the surrogate based optimization method</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>ego</td>
<td>Use the Efficient Global Optimization method</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>ea</td>
<td>Use an evolutionary algorithm</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>rng</td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>
Description

global_evidence allows the user to specify several global approaches for calculating the belief and plausibility functions:

- **lhs**: note: this takes the minimum and maximum of the samples as the bounds per "interval cell combination."
- **ego**: uses Efficient Global Optimization which is based on an adaptive Gaussian process surrogate.
- **sbo**: uses a Gaussian process surrogate (non-adaptive) within an optimization process.
- **ea**: uses an evolutionary algorithm. This can be expensive as the ea will be run for each interval cell combination.

Note that to calculate the plausibility and belief cumulative distribution functions, one has to look at all combinations of intervals for the uncertain variables. In terms of implementation, if one is using LHS sampling as outlined above, this method creates a large sample over the response surface, then examines each cell to determine the minimum and maximum sample values within each cell. To do this, one needs to set the number of samples relatively high: the default is 10,000 and we recommend at least that number. If the model you are running is a simulation that is computationally quite expensive, we recommend that you set up a surrogate model within the Dakota input file so that global_evidence performs its sampling and calculations on the surrogate and not on the original model. If one uses optimization methods instead to find the minimum and maximum sample values within each cell, this can also be computationally expensive.

Additional Resources

See the topic page evidence_theory for important background information and usage notes.

Refer to variable_support for information on supported variable types.

Theory

The basic idea is that one specifies an "evidence structure" on uncertain inputs and propagates that to obtain belief and plausibility functions on the response functions. The inputs are defined by sets of intervals and Basic Probability Assignments (BPAs). Evidence propagation is computationally expensive, since the minimum and maximum function value must be calculated for each "interval cell combination." These bounds are aggregated into belief and plausibility.

See Also

These keywords may also be of interest:

- **global_interval_est**
- **local_evidence**
- **local_interval_est**

7.2.66.1 samples

- **Keywords Area**
- **method**
- **global_evidence**
• **samples**

Number of samples for sampling-based methods

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

The `samples` keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

**Default Behavior**

By default, Dakota will use the minimum number of samples required by the chosen method.

**Usage Tips**

To obtain linear sensitivities or to construct a linear response surface, at least \( \text{dim} + 1 \) samples should be used, where \( \text{dim} \) is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \( (\text{dim} + 1)(\text{dim} + 2)/2 \) samples are needed. For uncertainty quantification, we recommend at least \( 10 \times \text{dim} \) samples. For `variance_based_decomp`, we recommend hundreds to thousands of samples. Note that for `variance_based_decomp`, the number of simulations performed will be \( N \times (\text{dim} + 2) \).

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
```

```
7.2.66.2  seed
```

• **Keywords Area**

• **method**

• **global_evidence**

• **seed**

Seed of the random number generator

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random `seed` control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.
Default Behavior
If not specified, the seed is randomly generated.

Expected Output
If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips
If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.66.3  sbo

- Keywords Area
- method
- global_evidence
- sbo

Use the surrogate based optimization method

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>gaussian_process</td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_-points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>
Description

A surrogate-based optimization method will be used. The surrogate employed in `sbo` is a Gaussian process surrogate.

The main difference between `ego` and the `sbo` approach is the objective function being optimized. `ego` relies on an expected improvement function, while in `sbo`, the optimization proceeds using an evolutionary algorithm (`coliny_ea`) on the Gaussian process surrogate: it is a standard surrogate-based optimization. Also note that the `sbo` option can support optimization over discrete variables (the discrete variables are relaxed) while `ego` cannot.

This is not the same as `surrogate_based_global`.

7.2.66.3.1 gaussian_process

- **Keywords Area**
  - method
  - global_evidence
  - sbo
  - gaussian_process

Gaussian Process surrogate model

Specification

**Alias:** kriging

**Argument(s):** none

**Default:** Surfpack Gaussian process

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Implementation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>surfpack</td>
<td>dakota</td>
<td>Use the Surfpack version of Gaussian Process surrogates Select the built in Gaussian Process surrogate</td>
</tr>
</tbody>
</table>

Description

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.
Use the Surfpack version of Gaussian Process surrogates

## Specification

**Alias:** none  
**Argument(s):** none

## Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. **Optimization methods:**
   - Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See [optimization_method](#).
   - The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. **Trend Function:**
   - The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See [trend](#).

3. **Correlation Lengths:**
   - Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See [correlation_lengths](#).

4. **Ill-conditioning**
   - One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   - Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   - The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using [surfpack](#).

5. **Gradient Enhanced Kriging (GEK).**
   - The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   - See notes in the Theory section.
Theory

**Gradient Enhanced Kriging**

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here “inexpensive” means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK’s correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging’s correlation matrix (which is a small sub-matrix within GEK’s correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK’s correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging’s correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK’s correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK’s correlation matrix.

7.2.66.3.1.2 dakota

- **Keywords Area**
- **method**
- **global_evidence**
- **sbo**
- **gaussian_process**
- **dakota**

Select the built in Gaussian Process surrogate

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

A second version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussion_process dakota model can be found in[60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the
selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

7.2.66.3.2 use_derivatives

- Keywords Area
- method
- global_evidence
- sbo
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.66.3.3 import_build_points_file

- Keywords Area
- method
- global_evidence
- sbo
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file
Argument(s): STRING
Default: no point import from a file
Child Keywords:
**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- `annotated` (default)
- `custom_annotated`
- `freeform`

**Examples**

```plaintext
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

**7.2.66.3.3.1 custom_annotated**

- **Keywords Area**
  - method
  - global_evidence
  - sbo
  - import_build_points_file
  - custom_annotated

Selects custom-annotated tabular file format
7.2 method

Topics

This keyword is related to the topics:

- file Formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.66.3.3.2 header

- Keywords Area
- method
- global_evidence
- sbo
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no header

**Description**

See description of parent `custom_annotated`

7.2.66.3.3.3 eval_id

- Keywords Area
- method
- global_evidence
- sbo
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
7.2 method

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.66.3.3.4 interface_id

• Keywords Area
• method
• global_evidence
• sbo
• import_build_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.66.3.3.5 annotated

• Keywords Area
• method
• global_evidence
• sbo
• import_build_points_file
• annotated

Selects annotated tabular file format
Topics

This keyword is related to the topics:

- `file_formats`

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```plaintext
%eval_id interface      x1    x2    obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID    0.9    1.1    0.0002    0.26    0.76
2   NO_ID    0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID    0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
7.2.6.6.3.6 freeform

• Keywords Area
  • method
  • global_evidence
  • sbo
  • import_build_points_file
  • freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996204857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.66.3.3.7 active_only

- Keywords Area
- method
- global_evidence
- sbo
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.66.3.4 export_approx_points_file

- Keywords Area
- method
- global_evidence
Output file for surrogate model value evaluations

### Specification

**Alias:** `export_points_file`  
**Argument(s):** STRING  
**Default:** no point export to a file  

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

### Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

### Usage Tips

Dakota exports tabular data in one of three formats:

- annotated *(default)*
- custom_annotated
- freeform

### custom_annotated

- **Keywords Area**
  - method
  - global_evidence
  - sbo
  - export_approx_points_file
  - custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.66.3.4.2 header

- Keywords Area
- method
- global_evidence
- sbo
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.66.3.4.3 eval_id

- Keywords Area
- method
- global_evidence
- sbo
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.66.3.4.4 interface_id

• Keywords Area
• method
• global_evidence
• sbo
• export_approx_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.66.3.4.5 annotated

• Keywords Area
• method
• global_evidence
• sbo
• export_approx_points_file
• annotated

Selects annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
7.2.66.3.4.6 freeform

- Keywords Area
- method
- global_evidence
- sbo
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1  0.0001996404857  0.2601620081  0.759955
0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.66.4  ego

- Keywords Area
- method
- global_evidence
- ego

Use the Efficient Global Optimization method

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>gaussian_process</td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_-points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>export_approx_-points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
</tbody>
</table>
Description

In the case of ego, the efficient global optimization (EGO) method is used to calculate bounds. By default, the Surfpack GP (Kriging) model is used, but the Dakota implementation may be selected instead. If use_derivatives is specified the GP model will be built using available derivative data (Surfpack GP only).

See efficient_global for more information.

7.2.66.4.1 gaussian_process

- Keywords Area
  - method
  - global_evidence
  - ego
  - gaussian_process

Gaussian Process surrogate model

Specification

Alias: kriging

Argument(s): none

Default: Surfpack Gaussian process

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group GP Implementation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dakota</td>
<td>Select the built in Gaussian Process surrogate</td>
</tr>
</tbody>
</table>

Description

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the surfpack keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. **In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.**

7.2.66.4.1.1 surfpack

- Keywords Area
  - method
Use the Surfpack version of Gaussian Process surrogates

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack. Several user options are available:

1. **Optimization methods:**
   
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default, the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See `optimization_method`.

   The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. **Trend Function:**
   
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See `trend`.

3. **Correlation Lengths:**
   
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See `correlation_lengths`.

4. **Ill-conditioning**
   
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.

   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.

   The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using

5. **Gradient Enhanced Kriging (GEK).**

   The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).

   See notes in the Theory section.
Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK’s correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging’s correlation matrix (which is a small sub-matrix within GEK’s correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK’s correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging’s correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK’s correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK’s correlation matrix.

7.2.66.4.1.2 dakota

- Keywords Area
- method
- global_evidence
- ego
- gaussian_process
- dakota

Select the built in Gaussian Process surrogate

Specification

Alias: none

Argument(s): none

Description

A second version of GP surrogates was available in prior versions of Dakota. For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in[60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the
selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

7.2.66.4.2 use_derivatives

- Keywords Area
- method
- global_evidence
- ego
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it’s use with Surfpack Gaussian process is not recommended.

7.2.66.4.3 import_build_points_file

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file

Argument(s): STRING

Default: no point import from a file

Child Keywords:
### Dakota Keyword

**Dakota Keyword**

- `custom_annotated`
  
  Selects custom-annotated tabular file format

- `annotated`
  
  Selects annotated tabular file format

- `freeform`
  
  Selects freeform file format

- `active_only`
  
  Import only active variables from tabular data file

### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- `custom_annotated`
- `freeform`

### Examples

```plaintext
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'

7.2.66.4.3.1 custom_annotated

- Keywords Area
- method
- global_evidence
- ego
- `import_build_points_file`
- `custom_annotated`

Selects custom-annotated tabular file format
```
7.2 method

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

```
%eval_id x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9   1.1  0.0002   0.26   0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.66.4.3.2  header

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.66.4.3.3  eval_id

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.66.4.3.4 interface_id

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.66.4.3.5 annotated

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- annotated

Selects annotated tabular file format
Topics
This keyword is related to the topics:

- file_formats

Specification
Alias: none
Argument(s): none
Default: annotated format

Description
An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior
By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples
Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID  0.9  1.1  0.0002  0.26  0.76
2   NO_ID  0.9009 1.1  0.0001996404857  0.2601620081  0.759955
3   NO_ID  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2.66.4.3.6 freeform

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
...```

7.2.66.4.3.7 active_only

- Keywords Area
- method
- global_evidence
- ego
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
7.2.6 method

- ego
- export_approx_points_file

Output file for surrogate model value evaluations

**Specification**

**Alias:** export_points_file

**Argument(s):** STRING

**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_ annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated *(default)*
- custom_ annotated
- freeform

7.2.6.4.4.1 custom_annotated

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- custom_ annotated

Selects custom-annotated tabular file format
Topics
This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom.annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
7.2 method

tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.66.4.4.2 header

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.66.4.4.3 eval_id

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent `custom_annotated`

7.2.66.4.4.4  interface_id

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent `custom_annotated`

7.2.66.4.4.5  annotated

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- annotated

Selects annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_f nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
7.2.66.4.4.6  freeform

- Keywords Area
- method
- global_evidence
- ego
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to **annotated** format, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In **freeform**, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.66.5 ea

- Keywords Area
- method
- global_evidence
- ea

Use an evolutionary algorithm

Specification

Alias: none
Argument(s): none

Description

In this approach, the evolutionary algorithm from Coliny, coliny_ea, is used to perform the interval optimization with no surrogate model involved. Again, this option of ea can support interval optimization over discrete variables.

7.2.66.6 lhs

- Keywords Area
- method
- global_evidence
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

Alias: none
Argument(s): none
Description

The lhs keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the lhs keyword must appear in conjunction with the sample_type keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

```
method
    sampling
        sample_type lhs
        samples = 20
```

7.2.66.7 response_levels

- Keywords Area
- method
- global_evidence
- response_levels

Values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response.levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>
Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

Default Behavior

If response_levels are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If response_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The num_response_levels is used to specify which arguments of the response_level correspond to which response.

Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

```plaintext
response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3
```

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num_response_levels key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```plaintext
method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05
variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = ’TF1n’ ’TF2n’
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = ’TF1u’ ’TF2u’
weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = ’TF1w’ ’TF2w’
histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
```
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.643154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844608686e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e+00</td>
<td>0.8000000000e+01</td>
<td></td>
</tr>
<tr>
<td>4.0400000000e+11</td>
<td>2.3000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.
In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.6.7.1 num_response_levels

- Keywords Area
- method
- global_evidence
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```
method sampling
  samples = 100
  seed = 34785
  num_response_levels = 1 1 1
  response_levels = 0.5 0.5 0.5
```

7.2.6.7.2 compute

- Keywords Area
- method
- global_evidence
- response_levels
• `compute`

Selection of statistics to compute at each response level

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td><code>probabilities</code></td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>gen_reliabilities</code></td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>system</code></td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

**Description**

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary_distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```
7.2 method

3.5e+05 4.0e+05 4.5e+05

compute reliabilities

7.2.66.7.2.1  probabilities

• Keywords Area
• method
• global_evidence
• response_levels
• compute
• probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The probabilities keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the probabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                  6.0e+04 6.5e+04 7.0e+04
                  3.5e+05 4.0e+05 4.5e+05
  compute probabilities

7.2.66.7.2.2  gen_reliabilities

• Keywords Area
• method
• global_evidence
• response_levels
• compute
• gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```
method
  sampling
    sample_type random
    samples = 100 seed = 1
complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

7.2.66.7.2.3 system

• Keywords Area
• method
• global_evidence
• response_levels
• compute
• system

Compute system reliability (series or parallel)

Specification

Alias: none
Argument(s): none
Child Keywords:
Description

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

### 7.2.66.7.2.4 series

- **Keywords Area**
- **method**
- **global_evidence**
- **response_levels**
- **compute**
- **system**
- **series**

**Specification**

- **Alias:** none
- **Argument(s):** none

**Description**

See parent keyword `system` for description.

### 7.2.66.7.2.5 parallel

- **Keywords Area**
- **method**
- **global_evidence**
- **response_levels**
• compute
• system
• parallel

Aggregate response statistics assuming a parallel system

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

See parent keyword `system` for description.

7.2.66.8  `probability_levels`

- Keywords Area
- method
- global_evidence
- `probability_levels`

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none  
**Argument(s):** `REALIST`  
**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not
very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```dakota
method,
  sampling,
  samples = 100 seed = 1
  complementary distribution
  probability_levels = 1.0 .66 .33 0.
    1.8 .5 0.
    1.3 .2 0.
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
  counts = 1 1
  descriptors = 'TF3h'
interface,
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.6000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of one result for each level mapping in the aleatory case).

7.2.66.8.1 num_probability_levels

- Keywords Area
- method
- global_evidence
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** probability_levels evenly distributed among response functions
7.2 method

Description

See parent page

7.2.66.9 gen_reliability_levels

- Keywords Area
- method
- global_evidence
- gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.66.9.1 num_gen_reliability_levels

- Keywords Area
- method
- global_evidence
• `gen_reliability_levels`
• `num_gen_reliability_levels`

Specify which `gen_reliability_levels` correspond to which response

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** `gen_reliability_levels` evenly distributed among response functions

**Description**

See parent page

7.2.66.10 distribution

• Keywords Area  
• method  
• `global_evidence`  
• `distribution`

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required/Optional Required <em>(Choose One)</em></td>
<td><code>cumulative</code></td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td><code>complementary</code></td>
<td></td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

**Description**

The `distribution` keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.
Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.66.10.1 cumulative

- Keywords Area
- method
- global_evidence
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

Alias: none

Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```
7.2.66.10.2 complementary

- Keywords Area
- method
- global_evidence
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
  distribution complementary
```

7.2.66.11 rng

- Keywords Area
- method
- global_evidence
- rng

Selection of a random number generator
7.2. method

Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>mt19937</td>
<td></td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td>rnum2</td>
<td></td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The **rng** keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the **rng** keyword must be accompanied by either **rnum2** (pseudo-random numbers) or **mt19937** (random numbers generated by the Mersenne twister). Otherwise, **mt19937**, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.66.11.1 mt19937

- Keywords Area
- method
- global_evidence
- rng
- mt19937

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none
Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on Wikipedia: http://en.wikipedia.org/wiki/Mersenne_twister.

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.66.11.2 `rnum2`

- Keywords Area
- method
- global_evidence
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package.

Specification

Alias: none

Argument(s): none

Description

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```
7.2 method

7.2.66.12 model_pointer

- Keywords Area
- method
- global_evidence
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ‘x1’ ‘x2’

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = ‘text_book’

responses
response_functions = 3
no_gradients
no_hessians

### 7.2.67 global_interval_est

- Keywords Area
- method
- global_interval_est

Interval analysis using global optimization methods

**Topics**

This keyword is related to the topics:

- uncertainty_quantification
- epistemic_uncertainty_quantification_methods
- interval_estimation

**Specification**

**Alias:** nond_global_interval_est

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Description

In the global approach to interval estimation, one uses either a global optimization method or a sampling method to assess the bounds of the responses.

`global_interval_est` allows the user to specify several approaches to calculate interval bounds on the output responses.

- **lhs** - note: this takes the minimum and maximum of the samples as the bounds
- **ego**
- **sbo**

<table>
<thead>
<tr>
<th>Optional (Choose One)</th>
<th>Solution Approach (Group 1)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>samples</strong></td>
<td>sbo</td>
<td>Use the surrogate based optimization method</td>
</tr>
<tr>
<td><strong>seed</strong></td>
<td>ego</td>
<td>Use the Efficient Global Optimization method</td>
</tr>
<tr>
<td><strong>max_iterations</strong></td>
<td>ea</td>
<td>Use an evolutionary algorithm</td>
</tr>
<tr>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td>rng</td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

Optional parameters include:

- **samples**: Number of samples for sampling-based methods
- **seed**: Seed of the random number generator
- **max_iterations**: Number of iterations allowed for optimizers and adaptive UQ methods
- **convergence_tolerance**: Stopping criterion based on objective function or statistics convergence
- **max_function_evaluations**: Number of function evaluations allowed for optimizers
- **rng**: Selection of a random number generator
- **model_pointer**: Identifier for model block to be used by a method
7.2.67.1 samples

Number of samples for sampling-based methods

Specification

**Alias:** none

**Argument(s):** INTEGER

**Description**

The `samples` keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

**Default Behavior**

By default, Dakota will use the minimum number of samples required by the chosen method.

**Usage Tips**

To obtain linear sensitivities or to construct a linear response surface, at least dim+1 samples should be used, where "dim" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least (dim+1)(dim+2)/2 samples are needed. For uncertainty quantification, we recommend at least 10*dim samples. For `variance_based_decomp`, we recommend hundreds to thousands of samples. Note that for `variance_based_decomp`, the number of simulations performed will be N*(dim+2).

**Examples**

```
method
  sampling
    sample_type lhs
    samples = 20
```
7.2.6.7.2 seed

- Keywords Area
- method
- global_interval_est
- seed

Seed of the random number generator

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.6.7.3 max_iterations

- Keywords Area
- method
- global_interval_est
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods
Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed $\text{max\_iterations}$ iterations. In cases where there are multiple nested iterations to be controlled, $\text{max\_iterations}$ generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations

7.2.67.4 convergence_tolerance

• Keywords Area
• method
• global_interval_est
• convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): REAL

Default: 1.e-4
Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do. Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.6 max_function_evaluations

- Keywords Area
- method
- global_interval_est
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 1000

Description

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.
Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

**Default Behavior**

Default value is 1000.

### 7.2.67.6 sbo

- **Keywords Area**
- **method**
- **global_interval_est**
- **sbo**

Use the surrogate based optimization method

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Optional</td>
<td></td>
<td>gaussian_process</td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
</tbody>
</table>

**Description**

A surrogate-based optimization method will be used. The surrogate employed in sbo is a Gaussian process surrogate.

The main difference between ego and the sbo approach is the objective function being optimized. ego relies on an expected improvement function, while in sbo, the optimization proceeds using an evolutionary algorithm (coliny_ea) on the Gaussian process surrogate: it is a standard surrogate-based optimization. Also note that the sbo option can support optimization over discrete variables (the discrete variables are relaxed) while ego cannot.

This is not the same as surrogate_based_global.
Gaussian Process surrogate model

Specification

**Alias:** kriging

**Argument(s):** none

**Default:** Surfpack Gaussian process

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group GP Implementation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
</tbody>
</table>

**Description**

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the `dakota` version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.
• surfpack

Use the Surfpack version of Gaussian Process surrogates

Specification

Alias: none

Argument(s): none

Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See optimization_method.
   The total number of evaluations of the likelihood function can be controlled using the max_trials keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See trend.

3. Correlation Lengths:
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See correlation_lengths.

4. Ill-conditioning
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   The surfpack model handles ill-conditioning internally by default, but behavior can be modified using

5. Gradient Enhanced Kriging (GEK).
   The use_derivatives keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   See notes in the Theory section.

Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here “inexpensive” means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use
derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.

7.2.67.6.1.2 dakota

• Keywords Area

• method

• global_interval_est

• sbo

• gaussian_process

• dakota

Select the built in Gaussian Process surrogate

Specification

Alias: none

Argument(s): none

Description

A second version of GP surrogates was available in prior versions of Dakota. For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota's. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in[60].

Dakota's GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a "nugget," but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.
Use the experimental Gaussian Process surrogate

**Specification**

**Alias**: none  
**Argument(s)**: none  

**Child Keywords**:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>optional</td>
<td></td>
<td>options_file</td>
<td>Filename for a YAML file that specifies Gaussian process options</td>
</tr>
</tbody>
</table>

**Description**

The experimental GP surrogate is an emerging capability intended to supersede the dakota and surfpack GP implementations by unifying the best features of both. It also includes additional features such as the ability to export the model and then use it from Python.

**See Also**

These keywords may also be of interest:

- experimental_gaussian_process
7.2 method

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no advanced options file

Description

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

Examples

```yaml
# FILE: gp_options.yaml

GP Parameters:
  Sigma Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
  scaler name: standardization
  num restarts: 20
Nugget:
  fixed nugget: 0.0
  estimate nugget: true
  Bounds:
    lower bound: 3.17e-8
    upper bound: 1.0e-2
  gp seed: 42
  standardize response: false
Trend:
  estimate trend: true
  Options:
    max degree: 2
    reduced basis: false
    p-norm: 1.0
    scaler type: none
    regression solver type: SVD
    verbosity: 1
kernel type: squared exponential
Length-scale Bounds:
  lower bound: 1.0e-2
  upper bound: 1.0e2
  verbosity: 1
```

7.2.67.6.2 **use_derivatives**

- **Keywords Area**
  - method
  - **global_interval_est**
  - sbo
  - **use_derivatives**

Use derivative data to construct surrogate models

Specification

**Alias:** none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.67.6.3 import_build_points_file

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file

Argument(s): STRING

Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

The import_build_points_file allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.
7.2 Method

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```plaintext
method
polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.67.6.3.1 custom_annotated

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tagular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
7.2.67.6.3.2 header

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no header

**Description**

See description of parent `custom_annotated`

7.2.67.6.3.3 eval_id

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`
7.2.67.6.3.4  interface_id

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

**Description**

See description of parent **custom_annotated**

7.2.67.6.3.5  annotated

- Keywords Area
- method
- global_interval_est
- sbo
- import_build_points_file
- annotated

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

- file Formats

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format
7.2 method

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  NO_ID   0.9  1.1  0.0002  0.26  0.76
2  NO_ID   0.9009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID   0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.67.6.3.6 freeform

- Keywords Area

- method

- global_interval_est

- sbo

- import_build_points_file

- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to **annotated** format, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In **freeform**, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment  
  tabular_data  
    tabular_data_file = 'dakota_summary.dat'  
    freeform
```

Resulting tabular file:

```
<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
...```

7.2.67.6.3.7 active_only

- **Keywords Area**
- **method**
- **global_interval_est**
- **sbo**
- **import_build_points_file**
- **active_only**

Import only active variables from tabular data file
7.2 method

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables. This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.67.6.4 export_approx_points_file

• Keywords Area

• method

• global_interval_est

• sbo

• export_approx_points_file

Output file for surrogate model value evaluations

Specification

Alias: export_points_file
Argument(s): STRING
Default: no point export to a file
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
</tr>
</tbody>
</table>
Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate's predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.67.6.4.1 custom_annotated

- Keywords Area
- method
- global_interval_est
- sbo
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   0.90009 1.1 0.0001996404857 0.2601620081 0.759955
2   0.89991 1.1 0.0002003604863 0.2598380081 0.760045
3    ...
```
7.2.67.6.4.2  header

- Keywords Area
- method
- global_interval_est
- sbo
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none
**Argument(s):** none
**Default:** no header

Description

See description of parent `custom_annotated`

7.2.67.6.4.3  eval_id

- Keywords Area
- method
- global_interval_est
- sbo
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

**Alias:** none
**Argument(s):** none
**Default:** no eval_id column

Description

See description of parent `custom_annotated`
7.2.6.4.4  interface_id

- Keywords Area
- method
- global_interval_est
- sbo
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.6.4.5  annotated

- Keywords Area
- method
- global_interval_est
- sbo
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Selects freeform file format

This keyword is related to the topics:

- file_formats
7.2 method

Specification

**Alias:** none
**Argument(s):** none
**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<p>| | | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
<td></td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
<td></td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.67.7 ego

- **Keywords Area**
- **method**
- **global_interval_est**
- **ego**

Use the Efficient Global Optimization method
Specification

**Alias:** none  
**Argument(s):** none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>gaussian_process</td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_-points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>export_approx_-points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
</tbody>
</table>

Description

In the case of `ego`, the efficient global optimization (EGO) method is used to calculate bounds. By default, the Surfpack GP (Kriging) model is used, but the Dakota implementation may be selected instead. If `use_derivatives` is specified the GP model will be built using available derivative data (Surfpack GP only).

See efficient_global for more information.

7.2.67.7.1 `gaussian_process`

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process

Gaussian Process surrogate model

Specification

**Alias:** kriging  
**Argument(s):** none

**Default:** Surfpack Gaussian process

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Group GP Implementation (Group 1)</td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dakota</td>
<td>Select the built in Gaussian Process surrogate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>experimental</td>
<td>Use the experimental Gaussian Process surrogate</td>
</tr>
</tbody>
</table>

### Description

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

#### 7.2.67.7.1.1 surfpack

- **Keywords Area**
  - method
  - global_interval_est
  - ego
  - gaussian_process
  - surfpack

Use the Surfpack version of Gaussian Process surrogates

### Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>export_model</td>
<td>export_model</td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
</tbody>
</table>
Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See `optimization_method`.
   The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See `trend`.

3. Correlation Lengths:
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See `correlation_lengths`.

4. Ill-conditioning
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using `ill_conditioning`.

5. Gradient Enhanced Kriging (GEK).
   The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   See notes in the Theory section.

Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.
7.2.67.1.2 export_model

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- surfpack
- export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>formats</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- text_archive - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- binary_archive - Binary, machine-readable archive for re-import or use with the surfpack executable
- algebraic_file - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- algebraic_console - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

Dakota Version 6.15 Reference Manual generated on November 5, 2021
These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior
No export.

Expected Output
Output depends on selected format; see the `formats` specification.

Additional Discussion
The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.67.7.1.3 filename_prefix

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- surfpack
- export_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics
This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
   algebraic_file
   binary_archive
```

Topics

This keyword is related to the topics:

- `surrogate_models`

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See `export_model` and the entries for the format selection keywords for further information.

#### 7.2.67.7.1.5 `text_archive`

- **Keywords Area**
- **method**
- **global_interval_est**
- **ego**
- **gaussian_process**
- **surfpack**
- **export_model**
- **formats**
- **text_archive**

Surrogate model plain-text archive file format

### Topics

This keyword is related to the topics:

- **surrogate_models**

### Specification

**Alias:** none  
**Argument(s):** none

### Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.
Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.6.7.1.6 binary_archive

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- surfpack
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which 'bsps' stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Select the built in Gaussian Process surrogate

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

A second version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.  

More details on the gaussian_process dakota model can be found in[60].  

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.  

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

**7.2.67.7.1.8 experimental**

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- experimental

Use the experimental Gaussian Process surrogate

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**
### Description

The experimental GP surrogate is an emerging capability intended to supersede the dakota and surfpack GP implementations by unifying the best features of both. It also includes additional features such as the ability to export the model and then use it from Python.

### See Also

These keywords may also be of interest:

- experimental_gaussian_process

### export_model

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- experimental
- export_model

Exports surrogate model in user-specified format(s)

### Topics

This keyword is related to the topics:

- surrogate_models

### Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:
### Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (`bin/surfpack`) or a user-developed tool. Export format is controlled using the `formats` specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive**: Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive**: Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file**: Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console**: Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

### Default Behavior

No export.

### Expected Output

Output depends on selected format; see the `formats` specification.

### Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
  formats
    algebraic_file
    binary_archive
```

7.2.67.7.1.11 formats

• Keywords Area
• method
• global_interval_est
• ego
• gaussian_process
Formats for surrogate model export

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>text_archive</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See export_model and the entries for the format selection keywords for further information.

7.2.67.7.1.12 text_archive

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- experimental
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format
7.2 method

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named {prefix}.{response_descriptor}.txt. Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.sps, in which 'sps' stands for Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.67.1.13 binary_archive

• Keywords Area
• method
• global_interval_est
• ego
• gaussian_process
• experimental
• export_model
• formats
• binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none
Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which ‘bsps’ stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.67.7.1.14 options_file

- Keywords Area
- method
- global_interval_est
- ego
- gaussian_process
- experimental
- options_file

Filename for a YAML file that specifies Gaussian process options

Specification

Alias: none

Argument(s): STRING

Default: no advanced options file

Description

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

Examples

# FILE: gp_options.yaml

GP Parameters:
  Sigma Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
  scaler name: standardization
  num restarts: 20
  Nugget:
    fixed nugget: 0.0
    estimate nugget: true
  Bounds:
    lower bound: 3.17e-8
7.2 method

upper bound: 1.0e-2
gp seed: 42
standardize response: false
Trend:
estimate trend: true
Options:
max degree: 2
reduced basis: false
p-norm: 1.0
scaler type: none
regression solver type: SVD
verbosity: 1
kernel type: squared exponential
Length-scale Bounds:
lower bound: 1.0e-2
upper bound: 1.0e2
verbosity: 1

7.2.67.7.2 use_derivatives

- Keywords Area
- method
- global_interval_est
- ego
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it's use with Surfpack Gaussian process is not recommended.

7.2.67.7.3 import_build_points_file

- Keywords Area
- method
- global_interval_est
- ego
- import_build_points_file

File containing points you wish to use to build a surrogate
Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional</strong> (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.67.7.3.1 custom_annotated

- **Keywords Area**
- **method**
Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.67.7.3.2 header

• Keywords Area
  • method
  • global_interval_est
  • ego
  • import_build_points_file
  • custom_annotated
  • header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

**See description of parent** custom_annotated

7.2.67.7.3.3 eval_id

• Keywords Area
  • method
  • global_interval_est
  • ego
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom.annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent `custom.annotated`
• **annotated**

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

• file_formats

**Specification**

*Alias:* none  
*Argument(s):* none  
*Default:* annotated format

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The *annotated* keyword can be used to explicitly specify this.

**Usage Tips**

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify *custom_annotated header eval_id*

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to *annotated format*, though *freeform* remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

**Examples**

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment  
tabular_data  
  tabular_data_file = 'dakota_summary.dat'  
  annotated
```

**Resulting tabular file:**

```
% eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

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7.2.6 freeform

- Keywords Area
- method
- global_interval_est
- ego
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.67.7.3.7 active_only

- Keywords Area
- method
- global_interval_est
- ego
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.67.7.4 export_approx_points_file

- Keywords Area
- method
- global_interval_est
**7.2 method**

- **ego**
- **export_approx_points_file**

Output file for surrogate model value evaluations

**Specification**

**Alias:** export_points_file

**Argument(s):** STRING

**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td><strong>Tabular Format</strong></td>
<td><strong>Group 1</strong></td>
<td><strong>custom_annotated</strong></td>
</tr>
<tr>
<td></td>
<td>annotated</td>
<td>annotated file format</td>
<td></td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>freeform file format</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- **annotated** (default)
- **custom_annotated**
- **freeform**

**7.2.67.7.4.1 custom_annotated**

- **Keywords Area**
  - **method**
  - **global_interval_est**
  - **ego**
  - **export_approx_points_file**
  - **custom_annotated**

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

• file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom.annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
7.2 method

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.67.7.4.2 header

- Keywords Area
- method
- global_interval_est
- ego
- export_approx_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

7.2.67.7.4.3 eval_id

- Keywords Area
- method
- global_interval_est
- ego
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent `custom_annotated`

7.2.67.7.4.4  **interface_id**

- Keywords Area
- method
- global_interval_est
- ego
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent `custom_annotated`

7.2.67.7.4.5  **annotated**

- Keywords Area
- method
- global_interval_est
- ego
- export_approx_points_file
- annotated

Selects annotated tabular file format
7.2 method

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface  x1   x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   NO_ID         0.9   1.1  0.0002  0.26              0.76
2   NO_ID  0.90009  1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID  0.89991  1.1 0.0002003604863 0.2598380081 0.760045
...```

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7.2.67.4.6 freeform

- Keywords Area
- method
- global_interval_est
- ego
- export_approx_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.9009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.67.8 ea

- Keywords Area
- method
- global_interval_est
- ea

Use an evolutionary algorithm

Specification

**Alias:** none

**Argument(s):** none

Description

In this approach, the evolutionary algorithm from Coliny, `coliny_ea`, is used to perform the interval optimization with no surrogate model involved. Again, this option of `ea` can support interval optimization over discrete variables.

7.2.67.9 lhs

- Keywords Area
- method
- global_interval_est
- lhs

Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

**Alias:** none

**Argument(s):** none
Description

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

```
method
  sampling
    sample_type lhs
    samples = 20
```

### 7.2.67.10  rng

- **Keywords Area**
- **method**
- **global_interval_est**
- **rng**

Selection of a random number generator

Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>RNG Algorithm (Group 1)</td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>
Description

The **rng** keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the **rng** keyword must be accompanied by either **rnum2** (pseudo-random numbers) or **mt19937** (random numbers generated by the Mersenne twister). Otherwise, **mt19937**, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

Generates random numbers using the Mersenne twister

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The **mt19937** keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

**mt19937** is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the **rng** keyword.

Usage Tips

Use of the Mersenne twister random number generator (**mt19937**) is recommended.

Examples

```
method
  sampling
    sample_type lhs
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
samples = 10
seed = 98765
rng mt19937

7.2.67.10.2 rnum2

• Keywords Area
• method
• global_interval_est
• rng
• rnum2

Generates pseudo-random numbers using the Pecos package

Specification

Alias: none

Argument(s): none

Description

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2

7.2.67.11 model_pointer

• Keywords Area
• method
• global_interval_est
• model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

• block_pointer
Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```dakota
environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
        response_levels = 0.1 0.2 0.6
        sample_type lhs
        distribution cumulative

model
    id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
    id_method = 'DACE'
    model_pointer = 'DACE_M'
    sampling sample_type lhs
        samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
    id_interface = 'I1'
    system asynch evaluation_concurrency = 5
```
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.68 bayes_calibration

- Keywords Area
- method
- bayes_calibration

Bayesian calibration

Topics

This keyword is related to the topics:

- bayesian_calibration
- package_queso

Specification

Alias: nond_bayes_calibration

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Bayesian Calibration Method (Group 1)</td>
<td>queso</td>
<td>Markov Chain Monte Carlo algorithms from the QUESO package</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gpmsa</td>
<td>(Experimental) Gaussian Process Models for Simulation Analysis (GPMSA) Bayesian calibration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wasabi</td>
<td>(Experimental Method) Non-MCMC Bayesian inference using interval analysis</td>
</tr>
<tr>
<td>Optional</td>
<td>experimental_design</td>
<td>(Experimental)</td>
<td>Adapatively select experimental designs for iterative Bayesian updating</td>
</tr>
<tr>
<td>Optional</td>
<td>calibrate_error_multipliers</td>
<td>Calibrate hyper-parameter multipliers on the observation error covariance</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>burn_in_samples</td>
<td>Manually specify the burn in period for the MCMC chain.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>posterior_stats</td>
<td>Compute information-theoretic metrics on posterior parameter distribution</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>chain_diagnostics</td>
<td>Compute diagnostic metrics for Markov chain</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_evidence</td>
<td>Calculate model evidence (marginal likelihood of model) when using Bayesian methods</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_discrepancy</td>
<td>(Experimental) Post-calibration calculation of model discrepancy correction</td>
<td></td>
</tr>
</tbody>
</table>

**7.2 method**

| dream | DREAM (DiffeRential Evolution Adaptive Metropolis) |
| muq | Markov Chain Monte Carlo algorithms from the MUQ package |
### Description

Bayesian calibration methods take prior information on parameter values (in the form of prior distributions) and observational data (e.g., from experiments) and infer posterior distributions on the parameter values. When the computational simulation is then executed with samples from the posterior parameter distributions, the results that are produced are consistent with (“agree with”) the experimental data. Calibrating parameters from a computational simulation model requires a likelihood function that specifies the likelihood of observing a particular observation given the model and its associated parameterization; Dakota assumes a Gaussian likelihood function. The algorithms that produce the posterior distributions on model parameters are most commonly Monte Carlo Markov Chain (MCMC) sampling algorithms. MCMC methods require many samples, often tens of thousands, so in the case of model calibration, often emulators of the computational simulation are used. For more details on the algorithms underlying the methods, see the Dakota User’s manual.

Dakota has four classes of Bayesian calibration methods: QUESO/DRAM, GPMSA, DREAM, and WASABI.

1. The QUESO methods use components from the QUESO library (Quantification of Uncertainty for Estimation, Simulation, and Optimization) developed at The University of Texas at Austin. Dakota uses its DRAM (Delayed Rejected Adaptive Metropolis) algorithm, and variants, for the MCMC sampling.

2. GPMSA (Gaussian Process Models for Simulation Analysis) is an approach developed at Los Alamos National Laboratory and Dakota currently uses the QUESO implementation. It constructs Gaussian process models to emulate the expensive computational simulation as well as model discrepancy. GPMSA also has extensive features for calibration, such as the capability to include a model discrepancy term and the capability to model functional data such as time series data. This is an experimental capability and not all features are available in Dakota yet.

3. DREAM (DiffeRential Evolution Adaptive Metropolis) is a method that runs multiple different chains simul-
automatically tunes the proposal covariance during the process by a self-adaptive randomized subspace sampling. [89].

4. WASABI: Non-MCMC Bayesian inference via interval analysis

**Usage Tips**

The Bayesian capabilities are under active development. At this stage, the QUESO methods in Dakota are the most advanced and robust, followed by DREAM, followed by GPMSA and WASABI which are not yet ready for production use.

The prior distribution is characterized by the properties of the uncertain variables. Correlated priors are only supported for unbounded normal, untruncated lognormal, uniform, exponential, gumbel, frechet, and weibull distributions and require specification of *standardized_space*, for example, for QUESO *standardized_space*

Note that as of Dakota 6.2, the field responses and associated field data may be used with QUESO and DREAM. That is, the user can specify field simulation data and field experiment data, and Dakota will interpolate and provide the proper residuals to the Bayesian calibration.

**Topics**

This keyword is related to the topics:

- bayesian_calibration
- package_queso

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Required</td>
<td>chain_samples</td>
<td>bayes_calibration</td>
<td>Number of Markov Chain Monte Carlo posterior samples</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td>Selection of a random number generator</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>emulator</td>
<td>Use an emulator or surrogate model to evaluate the likelihood function</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>standardized_\text{-}space</td>
<td>Perform Bayesian inference in standardized probability space</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>logit_transform</td>
<td>Utilize the logit transformation to reduce sample rejection for bounded domains</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_chainˌpointsˌfile</td>
<td>Export the MCMC chain to the specified filename</td>
<td></td>
</tr>
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</table>

**MCMC Algorithm (Group 1)**

<table>
<thead>
<tr>
<th>Optional/Choose One</th>
<th>dram</th>
<th>Use the DRAM MCMC algorithm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>delayed_rejection</td>
<td>Use the Delayed Rejection MCMC algorithm</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>adaptive_ˌmetropolis</td>
<td>Use the Adaptive Metropolis MCMC algorithm</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>metropolisˌhastings</td>
<td>Use the Metropolis-Hastings MCMC algorithm</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>multilevel</td>
<td>Use the multilevel MCMC algorithm</td>
</tr>
</tbody>
</table>

| Optional | pre_solve | Perform deterministic optimization for MAP before Bayesian calibration |
Description

The `queso` method supports the following MCMC algorithms: DRAM (Delayed Rejection Adaptive Metropolis), delayed rejection (DR) only, adaptive metropolis (AM) only, pure Metropolis Hasting (MH)s, and multilevel (ML).

When calibrating fast-running simulation models, use of an emulator is not typically warranted. For slower models, using an emulator model in the MCMC sampling will greatly improve the speed, since the Monte Carlo Markov Chain will generate thousands of samples on the emulator instead of the real simulation code. An emulator may be specified with the keyword `emulator`, followed by a `gaussian_process` emulator, a `pce` emulator (polynomial chaos expansion), or a `sc` emulator (stochastic collocation). For the `gaussian_process` emulator, the user must specify whether to use the `surfpack` or `dakota` version of the Gaussian process. The user can define the number of samples `build_samples` from which the emulator should be built. It is also possible to build the Gaussian process from points read in from the `import_points_file` and to export approximation-based sample evaluations using `export_points_file`. For `pce` or `sc`, the user can define a `sparse_grid_level`.

There are a variety of ways the user can specify the proposal covariance matrix which is very important in governing the samples generated in the chain. The proposal covariance specifies the covariance structure of a multivariate normal distribution. The user can specify `proposal_covariance`, followed by `derivatives`, `prior`, `values`, or `filename`. The derivative specification involves forming the Hessian of the misfit function (the negative log likelihood). When derivative information is available inexpensively (e.g. from an emulator), the derived-based proposal covariance forms a more accurate proposal distribution, resulting in lower rejection rates and faster chain mixing. The prior setting involves constructing the proposal from the variance of the prior distributions of the parameters being calibrated. When specifying the proposal covariance with values or from a file, the user can choose to specify only the diagonals of the covariance matrix with `diagonal` or to specify the full covariance matrix with `matrix`.

There are two other controls for QUESO. The `pre_solve` option enables the user to start the chain at an optimal point, the Maximum A Posteriori (MAP) point. This is the point in parameter space that maximizes the log posterior, (defined as the log-likelihood minus the log_prior). A deterministic optimization method is used to obtain the MAP point, and the MCMC chain is then started at the best point found in the optimization. The second factor is a `logit_transform`, which performs an internal variable transformation from bounded domains to unbounded domains in order to reduce sample rejection due to an out-of-bounds condition.

Note that as of Dakota 6.2, the field data capability may be used with QUESO. That is, the user can specify field simulation data and field experiment data, and Dakota will interpolate and provide the proper residuals to the Bayesian calibration.

7.2.68.1.1 chain_samples

- Keywords Area
- method
- bayes_calibration
- queso
- chain_samples

Number of Markov Chain Monte Carlo posterior samples
Specification

**Alias:** samples

**Argument(s):** INTEGER

**Default:** method-dependent

**Description**

The `chain_samples` keyword indicates the number of draws from the posterior distribution to perform. When an emulator is active, this will be the number of samples on the constructed surrogate model.

**Default Behavior**

The default number of chain samples is method-dependent. QUESO and MUQ methods use 1000. DREAM uses

(number of generations) x (number of chains), resulting in chain_samples close to that specified.

**Usage Tips**

MCMC methods typically require a large number of chain samples to converge, often thousands to millions.

**Examples**

```plaintext
method bayes_calibration queso
  chain_samples = 20000
```

7.2.68.1.2 seed

- Keywords Area
- method
- bayes_calibration
  - queso
- seed

Seed of the random number generator

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random `seed` control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.
Usage Tips

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.68.1.3 `rng`

- Keywords Area
- method
- bayes_calibration
- queso
- rng

Selection of a random number generator

Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

Child Keywords:

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

Description

The `rng` keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.
Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.68.1.3.1 mt19937

- Keywords Area
- method
- bayes_calibration
- queso
- rng
- mt19937

Generates random numbers using the Mersenne twister

Specification

**Alias:** none

**Argument(s):** none

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.68.1.3.2 rnum2

- Keywords Area
- method
- bayes_calibration
- queso
Generates pseudo-random numbers using the Pecos package

Specification

Alias: none

Argument(s): none

Description

The \texttt{rnum2} keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

\texttt{rnum2} is not used by default. To change this behavior, it must be specified in conjunction with the \texttt{rng} keyword.

Usage Tips

Use of the Mersenne twister random number generator (\texttt{mt19937}) is recommended over \texttt{rnum2}.

Examples

\begin{verbatim}
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
\end{verbatim}

7.2.68.1.4 emulator

• Keywords Area
• method
• bayes_calibration
• queso
• emulator

Use an emulator or surrogate model to evaluate the likelihood function

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

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### Description

This keyword describes the type of emulator used when calculating the likelihood function for the Bayesian calibration. The emulator can be a Gaussian process, polynomial chaos expansion, or stochastic collocation.

#### 7.2.68.1.4.1 gaussian_process

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process

Gaussian Process surrogate model

### Specification

**Alias:** kriging

**Argument(s):** none

**Default:** Surfpack Gaussian process

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required (Choose One)</th>
<th>Emulator Type (Group 1)</th>
<th>Surrogate Model</th>
</tr>
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<tbody>
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<td></td>
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<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>pce</td>
<td>Polynomial Chaos Expansion surrogate model</td>
</tr>
<tr>
<td></td>
<td>ml_pce</td>
<td>Multilevel Polynomial Chaos Expansion as an emulator model.</td>
</tr>
<tr>
<td></td>
<td>mf_pce</td>
<td>Multifidelity Polynomial Chaos Expansion as an emulator model.</td>
</tr>
<tr>
<td></td>
<td>sc</td>
<td>Stochastic Collocation as an emulator model.</td>
</tr>
<tr>
<td></td>
<td>mf_sc</td>
<td>Multifidelity Stochastic Collocation as an emulator model.</td>
</tr>
</tbody>
</table>
Description

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the `dakota` version is deprecated and intended to be removed in a future release.**

*Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.*

### 7.2.68.1.4.2 surfpack

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **gaussian_process**
- **surfpack**

Use the Surfpack version of Gaussian Process surrogates

#### Specification

**Alias:** none  
**Argument(s):** none
Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   - Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See optimization_method.
   - The total number of evaluations of the likelihood function can be controlled using the max_trials keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   - The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See trend.

3. Correlation Lengths:
   - Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See correlation_lengths.

4. Ill-conditioning
   - One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   - Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   - The surfpack model handles ill-conditioning internally by default, but behavior can be modified using 5. Gradient Enhanced Kriging (GEK).

4.5. Gradient Enhanced Kriging (GEK)
   - The use_derivatives keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   - See notes in the Theory section.

Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.
Select the built in Gaussian Process surrogate

**Specification**

- **Alias:** none
- **Argument(s):** none

**Description**

A second version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the [gaussian_process dakota model](#) can be found in [60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a **point_selection** option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the **point_selection** option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.
Specification

Alias: none
Argument(s): INTEGER

Description

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of build_samples is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

Examples

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```plaintext
method
  gpais
    build_samples = 100
    samples_on_emulator = 100
    max_iterations = 5
    response_levels = -1.065
```

7.2.68.1.4.5 posterior_adaptive

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- posterior_adaptive

Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none

Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.
Examples

bayes_calibration queso
    chain_samples = 2000 seed = 348
delayed_rejection
emulator
gaussian_process surfpack build_samples = 30
posterior_adaptive max_iterations = 10
proposal_covariance derivatives

7.2.68.1.4.6 import_build_points_file

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file

Argument(s): STRING

Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

active_only

Import only active variables from tabular data file

Description

The import_build_points_file allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.
Default Behavior
By default, methods do not import points from a file.

Usage Tips
Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = "dakota_uq_rosenbrock_pce_import.annot.pts.dat"
```

7.2.68.1.4.7 custom_annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```environment
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
```

7.2.68.1.4.8 header

• Keywords Area

• method
Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no header

**Description**

See description of parent `custom_annotated`

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`
7.2.68.1.4.10  interface_id

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.1.4.11  annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
    environment
    tabular_data
      tabular_data_file = 'dakota_summary.dat'
      annotated
```

Resulting tabular file:

```
#eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID   0.9 1.1 0.0002 0.26 0.76
2   NO_ID   0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.1.4.12  freeform

- Keywords Area
  - method
  - bayes_calibration
  - queso
  - emulator
Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```dakota_version

environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform

Resulting tabular file:

       0.9     1.1  0.0002  0.26   0.76
0.90009 1.1  0.0001996404857 0.2601620081 0.759955
0.89991 1.1  0.0002003604863 0.2598380081 0.760045
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
7.2.68.1.4.13  active_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- gaussian_process
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics
This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description
By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.14  pce

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce

Polynomial Chaos Expansion surrogate model
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
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<td>Group 1</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td></td>
<td></td>
<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cubature_integrand</td>
<td>Cubature using Stroud rules and their extensions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_order</td>
<td>The (initial) order of a polynomial expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
</tbody>
</table>
## Description

Selects a polynomial chaos expansion (PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using PCE as a surrogate within the Bayesian framework.

## See Also

These keywords may also be of interest:

- `polynomial_chaos`

7.2.68.1.4.15  `p_refinement`

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
- `emulator`
- `pce`
- `p_refinement`

Automatic polynomial order refinement
Specification

Alias: none
Argument(s): none
Default: no refinement
Child Keywords:

<table>
<thead>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>p-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.</td>
</tr>
</tbody>
</table>

Description

The p_refinement keyword specifies the usage of automated polynomial order refinement, which can be either uniform or dimension_adaptive.

The dimension_adaptive option is supported for the tensor-product quadrature and Smolyak sparse grid options and uniform is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the max_iterations and convergence_tolerance method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The dimension_adaptive case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the uniform refinement case, the quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
7.2.68.1.4.16  uniform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.1.4.17  dimension_adaptive

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.1.4.18 sobol

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.
Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```
method, polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```

7.2.68.1.4.19 decay

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• p_refinement
• dimension_adaptive
• decay

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

Specification

Alias: none
Argument(s): none

Description

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.
Examples

method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive p_refinement decay
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.1.4.20 generalized

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- p_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none

Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.1.4.21 max_refinement_iterations

- Keywords Area
- method
- bayes_calibration
7.2 method

- queso
- emulator
- pce
- max_refinement_iterations

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.

**Examples**

```plaintext
method,
polynomial_chaos
  dimension_adaptive generalized
  p_refinement
    max_refinement_iterations = 20
    convergence_tol = 1.e-4
    sparse_grid_level = 1
```

7.2.68.1.4.22 quadrature_order

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**
Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with `quadrature_order` and, optionally, `dimension_preference`). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a `dimension_preference`, where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified `quadrature_order` and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n} (p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the `quadrature_order` specification supports an array input.

A corresponding sequence specification is documented at, e.g., `quadrature_order_sequence` and `quadrature_order_sequence_sequence`.

**See Also**

These keywords may also be of interest:

- `quadrature_order_sequence` and `method-multifidelity_stoch_collocation-quadrature_order_sequence`

7.2.68.1.4.23 `dimension_preference`

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
7.2 method

- emulator
- pce
- quadrature_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.14.24 nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- quadrature_order
- nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested
Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.25  non_nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.1.4.26  sparse_grid_level

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

Alias: none
Argument(s): INTEGER
Child Keywords:
Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with sparse_grid_level and, optionally, dimension_preference). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the dimension_preference specification enables the use of anisotropic sparse grids (refer to the PCE description in the User's Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full sparse_grid_level and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User's Manual for additional details). As for quadrature_order, the sparse_grid_level specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., sparse_grid_level_sequence and sparse_grid_level_sequence

See Also

These keywords may also be of interest:

- sparse_grid_level_sequence method-multifidelity_stoch_collocation-sparse_grid_level_sequence
7.2.68.1.4.27  dimension_preference

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- sparse_grid_level
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leada to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.1.4.28  restricted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.
7.2 method

Specification

Alias: none

Argument(s): none

Default: restricted (except for generalized sparse grids)

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of $m = 2l + 1$ are used to enforce odd quadrature orders, where $l$ is the grid level and $m$ is the number of points in the rule. The precision of this Gauss rule is then $i = 2m - 1 = 4l + 1$. For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level $l$ for a sparse grid ($i = 4l + 1$) or an order $m$ for a tensor grid ($i = 2m - 1$). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.1.4.29 unrestricted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of $m = 2l + 1$ are used to enforce odd quadrature orders, where $l$ is the grid level and $m$ is the number of points in the rule. The precision of this Gauss rule is then $i = 2m - 1 = 4l + 1$. For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level $l$ for a sparse grid ($i = 4l + 1$) or an order $m$ for a tensor grid ($i = 2m - 1$). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.1.4.30 nested
   • Keywords Area
   • method
   • bayes_calibration
   • queso
   • emulator
   • pce
   • sparse_grid_level
   • nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.31 non_nested
   • Keywords Area
   • method
   • bayes_calibration
   • queso
   • emulator
   • pce
   • sparse_grid_level
   • non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none

Argument(s): none
Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.1.4.32  cubature_integrand

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- cubature_integrand

Cubature using Stroud rules and their extensions.

Specification

**Alias:** none

**Argument(s):** INTEGER

Description

Multi-dimensional integration by Stroud cubature rules [80] and extensions [94], as specified with cubature_integrand. A total-order expansion is used, where the isotropic order $p$ of the expansion is half of the integrand order, rounded down. The total number of terms $N$ for an isotropic total-order expansion of order $p$ over $n$ variables is given by

$$N = 1 + P = 1 + \sum_{s=1}^{P} \frac{1}{s!} \prod_{r=0}^{s-1} (n + r) = \frac{(n + p)!}{n!p!}$$

Since the maximum integrand order is currently five for normal and uniform and two for all other types, at most second- and first-order expansions, respectively, will be used. As a result, cubature is primarily useful for global sensitivity analysis, where the Sobol' indices will provide main effects and, at most, two-way interactions. In addition, the random variable set must be independent and identically distributed (iid), so the use of askey or wiener transformations may be required to create iid variable sets in the transformed space (as well as to allow usage of the higher order cubature rules for normal and uniform). Note that global sensitivity analysis often assumes uniform bounded regions, rather than precise probability distributions, so the iid restriction would not be problematic in that case.

7.2.68.1.4.33  expansion_order

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order

The (initial) order of a polynomial expansion.
Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

<table>
<thead>
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<th>Required/Optional</th>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>collocation_points</td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td></td>
<td></td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_samples</td>
<td>Number of simulation samples used to estimate the expected value of a set of PCE coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>posterior_adaptive</td>
<td>Adapt emulator model to increase accuracy in high posterior probability regions</td>
</tr>
</tbody>
</table>

Description

When the expansion_order for a polynomial chaos expansion is specified, the coefficients may be computed by integration based on random samples or by regression using either random or sub-sampled tensor product
7.2 method

quadrature points.

Multidimensional integration by Latin hypercube sampling (specified with expansion_samples). In this case, the expansion order $p$ cannot be inferred from the numerical integration specification and it is necessary to provide an expansion_order to specify $p$ for a total-order expansion.

Linear regression (specified with either collocation_points or collocation_ratio). A total-order expansion is used and must be specified using expansion_order as described in the previous option. To avoid requiring the user to calculate $N$ from $n$ and $p$, the collocation_ratio allows for specification of a constant factor applied to $N$ (e.g., collocation_ratio = 2 produces samples = $2N$). In addition, the default linear relationship with $N$ can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $cN_{pT}$). When admissible, a constrained least squares approach is employed in which response values are first reproduced exactly and error in reproducing response derivatives is minimized. Two collocation grid options are supported: the default is Latin hypercube sampling ("point collocation"), and an alternate approach of "probabilistic collocation" is also available through inclusion of the tensor_grid keyword. In this alternate case, the collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

If collocation_points or collocation_ratio is specified, the PCE coefficients will be determined by regression. If no regression specification is provided, appropriate defaults are defined. Specifically SVD-based least-squares will be used for solving over-determined systems and under-determined systems will be solved using LASSO. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares. Technical information on the various methods listed below can be found in the Linear regression section of the Theory Manual. Some of the regression methods (OMP, LASSO, and LARS) are able to produce a set of possible PCE coefficient vectors (see the Linear regression section in the Theory Manual). If cross validation is inactive, then only one solution, consistent with the noise_tolerance, will be returned. If cross validation is active, Dakota will choose between possible coefficient vectors found internally by the regression method across the set of expansion orders (1,...,expansion_order) and the set of specified noise tolerances and return the one with the lowest cross validation error indicator.

7.2.68.1.4.34 dimension_preference

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none
Argument(s): REALLIST
Default: isotropic grids
Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

• sobol
• decay

7.2.68.1.4.35 basis_type

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• expansion_order
• basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
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<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
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</table>
7.2 method

| adapted | Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. |

Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.

7.2.68.1.4.36 tensor_product

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- basis_type
- tensor_product

Use a tensor-product index set to construct a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Description

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$

7.2.68.1.4.37 total_order

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$
\sum_{k=1}^{d} i_k \leq p
$$

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- basis_type
- adapted
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<tr>
<td>Optional</td>
<td>Description of Group</td>
<td>advancements</td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td>Description of Group</td>
<td>soft_convergence_limit</td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.1.4.39 advancements

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **pce**
- **expansion_order**
- **basis_type**
- **adapted**
- **advancements**

The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.1.4.40  soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.

Specification

Alias: none

Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.1.4.41  collocation_points

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points

Number of collocation points used to estimate expansion coefficients

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:
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<th>Description of Group</th>
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<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
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<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
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<td></td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
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<td></td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>least_angle_regression</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
</tbody>
</table>
### Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.  
A corresponding sequence specification is documented at, e.g., `collocation_points_sequence` and `collocation_points_sequence`.

### See Also

These keywords may also be of interest:

- `collocation_points_sequence` method-multilevel_polynomial_chaos-expansion_order_sequence-collocation-
7.2 method

7.2.68.1.4.42 least_squares

- Keywords Area
- method
- bayes_calibration
- qeso
- emulator
- pce
- expansion_order
- collocation_points
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

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<tr>
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<th>Description of Group LSQ Regression Approach (Group 1)</th>
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<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
<td>svd</td>
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</tr>
</thead>
<tbody>
<tr>
<td>equality_-constrained</td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.
Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2 Method 2127

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.1.4.45 orthogonal_matching_pursuit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none

Child Keywords:

| Required/Optional | Description of Group | Dakota Keyword | Dakota Keyword
|-------------------|-----------------------|----------------|-----------------------|
| Optional          | noise_tolerance       | noise_tolerance| The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.1.4.46 noise_tolerance

- Keywords Area
- method
- bayes_calibration
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.47 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

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<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.1.4.49 noise_tolerance
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALIST

**Default:** $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.50 least_angle_regression

- **Keywords Area**
  - method
  - bayes_calibration
  - quo\(e\)
  - emulator
  - pce
  - expansion_order
  - collocation_points
  - least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

**Specification**

**Alias:** lars

**Argument(s):** none

**Child Keywords:**

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<thead>
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<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
7.2.68.1.4.51 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.52 least_absolute_shrinkage

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso

**Argument(s):** none

**Child Keywords:**

---

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</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td>l2_penalty</td>
<td>The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

### Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.1.4.53 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
7.2.68.1.4.54  l2_penalty

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.

7.2.68.1.4.55  cross_validation

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.1.4.56 noise_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.

Specification

Alias: none
Argument(s): none
Default: false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.
7.2.68.1.4.57  max_cv_order_candidates

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

 Specification

 **Alias:** none

 **Argument(s):** INTEGER

 **Default:** ushort max

 **Description**

 When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

 **See Also**

 These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train
Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

### Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

### Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the collocation_ratio allows for specification of a constant factor applied to \( N \) (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^{o} \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order.

### Perform bounds-scaling on response values prior to surrogate emulation
7.2 method

Specification

Alias: none
Argument(s): none
Default: use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to \([0,1]\) based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- multifidelity_polynomial_ chaos
- multifidelity_function_train

7.2.68.1.4.60 use_derivatives

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.
7.2.68.1.4.61  tensor_grid

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.1.4.62  reuse_points

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
Specify

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

`7.2.68.1.4.63 max_solver_iterations`

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_points
- max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specify

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

`7.2.68.1.4.64 collocation_ratio`

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

**Specification**

**Alias:** none  
**Argument(s):** REAL

**Child Keywords:**

<table>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
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<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
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<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
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<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td>Method</td>
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<td></td>
</tr>
<tr>
<td>-------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>least_angle_regression</code></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>least_absolute_shrinkage</code></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>cross_validation</strong></td>
<td>Use cross validation to choose the ‘best’ polynomial order of a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>ratio_order</strong></td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>response_scaling</strong></td>
<td>Perform bounds-scaling on response values prior to surrogate emulation.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>use_derivatives</strong></td>
<td>Use derivative data to construct surrogate models.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>tensor_grid</strong></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong> <strong>reuse_points</strong></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>-----------------------</td>
<td>----------------------------------------------------------</td>
<td></td>
</tr>
</tbody>
</table>

### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the collocation_ratio allows for specification of a constant factor applied to \( N \) (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^{o+1} \)).

7.2.68.1.4.65 least_squares

- **Keywords Area**
  - method
  - bayes_calibration
  - queso
  - emulator
  - pce
  - expansion_order
  - collocation_ratio
  - least_squares

Compute the coefficients of a polynomial expansion using least squares

### Specification

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group LSQ Regression Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td><strong>Optional/Choose One</strong></td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
<td>svd</td>
<td>svd</td>
</tr>
</tbody>
</table>
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Description**

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.68.1.4.66  **svd**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_ratio**
- **least_squares**
- **svd**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

7.2.68.1.4.67  **equality_constrained**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

### Specification

**Alias:** none  
**Argument(s):** none

### Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.1.4.68 orthogonal_matching_pursuit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

### Specification

**Alias:** omp  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.1.4.69 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

**Alias**: none

**Argument(s)**: REALLIST

**Default**: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.70 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Specification

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.1.4.71 basis_pursuit_denoising

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
| Optional          | noise_tolerance       |                | The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.
7.2.68.1.4.72 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.73 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:
### Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.1.4.74  noise_tolerance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_ratio**
- **least_angle_regression**
- **noise_tolerance**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### Specification

**Alias:** none

**Argument(s):** REALIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.75  least_absolute_shrinkage

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **pce**
7.2 method

- expansion_order
- collocation_ratio
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td>The l2 penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.1.4.76 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.77 l2_penalty

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

Specification

Alias: none

Argument(s): REAL

Default: 0. (reverts to standard LASSO formulation)

Description

The $l_2$ penalty used when performing compressed sensing with elastic net.

7.2.68.1.4.78 cross_validation

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

### Specification

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

### Description

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.1.4.79  noise_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.
Specification

Alias: none
Argument(s): none
Default: false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.1.4.80 max_cv_order_candidates

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.
Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.

**Description**

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \(cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
7.2.68.1.4.82  response_scaling

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• expansion_order
• collocation_ratio
• response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

Specification

Alias: none
Argument(s): none
Default: use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to \([0,1]\) based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

• multifidelity_polynomial_chaos
• multifidelity_function_train

7.2.68.1.4.83  use_derivatives

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
7.2 method

- expansion_order
- collocation_ratio
- use_derivatives

Use derivative data to construct surrogate models

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.1.4.84 tensor_grid

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
7.2.68.1.4.85  reuse_points

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.1.4.86  max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients
7.2 method

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.1.4.87 expansion_samples

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
  - expansion_order
- expansion_samples

Number of simulation samples used to estimate the expected value of a set of PCE coefficients

Specification

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

Description

In this case, the expansion order $p$ cannot be inferred from the numerical integration specification and it is necessary to provide an expansion_order to specify $p$ for a total-order expansion.

A corresponding sequence specification is documented at, e.g., expansion_samples_sequence and expansion_samples_sequence
See Also

These keywords may also be of interest:

- expansion_samples_sequence
- method
- multilevel_polynomial_chaos
- expansion_order_sequence
- expansion_samples_sequence

7.2.68.1.4.88 reuse_points

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- expansion_samples
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.68.1.4.89 import_build_points_file

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- import_build_points_file

File containing points you wish to use to build a surrogate
7.2.6 method

Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```
methode
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.1.4.90 custom_annotated

- Keywords Area
- method
**Selects custom-annotated tabular file format**

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.
Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   0.9  1.1  0.0002  0.26       0.76
2  0.90009  1.1 0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1 0.0002003604863  0.2598380081  0.760045
...```

7.2.68.1.4.91 header

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- expansion_order
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.1.4.94  annotated

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• expansion_order
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom.annotated` header `eval_id`.

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID  0.9  1.1  0.0002  0.26  0.76
2   NO_ID  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3   NO_ID  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
... 
```

7.2.68.1.4.95 freeform

• Keywords Area

• method

• bayes_calibration

• queso

• emulator

• pce

• expansion_order

• import_build_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9  1.1   0.0002  0.26  0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.1.4.96 active_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
• expansion_order
• import_build_points_file
• active_only

Import only active variables from tabular data file

Topics
This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description
By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.97 posterior_adaptive

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• expansion_order
• posterior_adaptive

Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none
7.2 method

Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

Examples

bayes_calibration queso
  chain_samples = 2000 seed = 348
delayed_rejection
delayed_rejection
  emulator
gaussian_process surfpack build_samples = 30
  posterior_adaptive max_iterations = 10
  proposal_covariance derivatives

7.2.68.1.4.98 orthogonal_least_interpolation

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

**Alias:** least_interpolation oli

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>collocation_points</td>
<td></td>
<td>Number of collocation points used to estimate expansion coefficients.</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>
Optional reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models. File containing points you wish to use to build a surrogate

Optional import_build_points_file

Optional posterior_adaptive

Adapt emulator model to increase accuracy in high posterior probability regions

Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods expansion_order cannot be set. OLI will produce the lowest degree polynomial that interpolates the data

7.2.68.1.4.99 collocation_points

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• orthogonal_least_interpolation
• collocation_points

Number of collocation points used to estimate expansion coefficients

Specification

Alias: none
Argument(s): INTEGER

Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.

A corresponding sequence specification is documented at, e.g., collocation_points_sequence and collocation_points_sequence
See Also

These keywords may also be of interest:

- collocation_points_sequence
- method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence

7.2.68.1.4.100 tensor_grid

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.1.4.101 reuse_points

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
### Specification

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

### Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

#### 7.2.68.1.4.102 import_build_points_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

### Specification

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>
Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.1.4.103 custom_annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias**: none

**Argument(s)**: none

**Default**: annotated format

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To control which header row and columns are in the input/output, specify **custom_annotated**, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to **annotated format**, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:
7.2 method

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.68.1.4.104 header

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.68.1.4.105 eval_id

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.68.1.4.106  interface_id

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• orthogonal_least_interpolation
• import_build_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.1.4.107  annotated

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• pce
• orthogonal_least_interpolation
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nlneq_con_1 nlneq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
7.2.68.1.4.108  freeform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to **annotated** format, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In **freeform**, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
```

7.2.68.1.4.109  active_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias**: none

**Argument(s)**: none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
Adapt emulator model to increase accuracy in high posterior probability regions

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

**Examples**

```plaintext
bayes_calibration queso
  chain_samples = 2000 seed = 348
  delayed_rejection
  emulator
    gaussian_process surfpack build_samples = 30
    posterior_adaptive max_iterations = 10
    proposal_covariance derivatives
```

**askey**

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.1.4.113  normalized

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

Specification

**Alias**: none

**Argument(s)**: none

**Default**: PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.1.4.114  export Expansion File

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

**Alias**: none

**Argument(s)**: STRING
7.2 method

Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.1.4.115  diagonal_covariance

• Keywords Area
  • method
  • bayes_calibration
  • queso
  • emulator
  • pce
  • diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none
Argument(s): none
Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.1.4.116  full_covariance

• Keywords Area
  • method
  • bayes_calibration
  • queso
  • emulator
  • pce
  • full_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none
Description

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.68.1.4.117  ml_pce

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce

Multilevel Polynomial Chaos Expansion as an emulator model.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td>Sample allocation approach for multilevel expansions</td>
</tr>
<tr>
<td>Optional</td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One) Group 1</td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askey</td>
</tr>
<tr>
<td>----------</td>
<td>-----------------------------------</td>
<td>------</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

<table>
<thead>
<tr>
<th>Optional</th>
<th>wiener</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

<table>
<thead>
<tr>
<th>Optional</th>
<th>normalized</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

<table>
<thead>
<tr>
<th>Optional</th>
<th>Covariance Type (Group 3)</th>
<th>diagonal_covariance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Display only the diagonal terms of the covariance matrix.

<table>
<thead>
<tr>
<th></th>
<th>full_covariance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Display the full covariance matrix.

**Description**

Selects a multilevel polynomial chaos expansion (ML PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using ML PCE as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- multilevel_polynomial_chaos

7.2.68.1.4.118 allocation_control

- Keywords Area
Sample allocation approach for multilevel expansions

**Specification**

*Alias*: none

*Argument(s):* none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Multilevel Sample Allocation Control (Group 1)</td>
<td>estimator_variance</td>
<td>Variance of mean estimator within multilevel polynomial chaos</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rip_sampling</td>
<td>Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

**Description**

Multilevel expansions, including regression-based polynomial chaos expansion (PCE) and function train (FT) expansions, require a sample allocation strategy. Three options are currently available:

- allocation based on assuming a convergence rate for the estimator variance (for regression PCE)
- restricted isometry property (RIP) sampling (for regression PCE via compressed sensing)
- rank sampling (for FT)

**Default Behavior**

Current defaults, when *allocation_control* is not specified, are estimator variance for PCE and rank sampling for FT.

7.2.68.1.4.119 estimator_variance
Variance of mean estimator within multilevel polynomial chaos

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>estimator_rate</td>
<td>Rate of convergence of mean estimator variance within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

**Description**

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}
\]

for free parameters \(\gamma\) and \(\kappa\), with default values that may be overridden as part of this specification block.

This approach is supported for regression-based PCE approaches (over-determined least squares, under-determined compressed sensing, or orthogonal least interpolation).

In practice, it can be challenging to estimate a smooth convergence rate for estimator variance in the presence of abrupt transitions in the quality of sparse recoveries. As a result, sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the convergence of the estimator variance, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.
method,
  model_pointer = 'HIERARCH'
  multilevel_polynomial_chaos
    orthogonal_matching_pursuit
    expansion_order_sequence = 2
    pilot_samples = 10
    collocation_ratio = .9
    allocation_control
    estimator_variance estimator_rate = 2.5
    seed = 1237
    convergence_tolerance = .01

7.2.68.1.4.120 estimator_rate

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- allocation_control
- estimator_variance
- estimator_rate

Rate of convergence of mean estimator within multilevel polynomial chaos

Speciﬁcation

Alias: none
Argument(s): REAL
Default: 2

Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[ \text{Var}[\hat{Q}] = \frac{\sigma_{\hat{Q}}^2}{N} \]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[ \text{Var}[\hat{Q}] = \frac{\sigma_{\hat{Q}}^2}{\gamma N^\kappa} \]

for free parameters \( \gamma \) and \( \kappa \).

The default values are \( \gamma = 1 \) and \( \kappa = 2 \) (adopts a more aggressive sample profile by assuming a faster convergence rate than Monte Carlo). This advanced speciﬁcation option allows to user to specify \( \kappa \), overriding the default.
Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos

**Specification**

**Alias:** none  
**Argument(s):** none  

**Description**

Multilevel polynomial chaos with compressed sensing may allocate the number of samples per level based on the restricted isometry property (RIP), applied to recovery at level $l$:

$$N_l \geq s_l \log^3(s_l) L_l \log(C_l)$$

for sparsity $s$, cardinality $C$, and mutual coherence $L$. The adaptive algorithm starts from a pilot sample, shapes the profile based on observed sparsity, and iterates until convergence. In practice, RIP sampling levels are quite conservative, and a collocation ratio constraint ($N_l \leq rC_l$, where $r$ defaults to 2) must be enforced on the profile.

The algorithm relies on observed sparsity, it is appropriate for use with regularized solvers for compressed sensing. It employs orthogonal matching pursuit (OMP) by default and automatically activates cross-validation in order to choose the best noise parameter value for the recovery.

This capability is **experimental**. Sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the number of sparse coefficient sets recovered for each level, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```dakota
method,  
model_pointer = 'HIERARCH'  
multilevel_polynomial_choas  
  orthogonal_matching_pursuit  
  expansion_order_sequence = 2  
  pilot_samples = 10  
  collocation_ratio = 0.9  
  allocation_control rip_sampling  
  seed = 1237  
  convergence_tolerance = 0.01
```
Formulation for emulation of model discrepancies.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** distinct  

**Child Keywords:**

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<tr>
<th>Required/Optional Description of Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required <em>(Choose One)</em> Discrepancy Emulation Approach (Group 1)</td>
<td>distinct</td>
</tr>
<tr>
<td>recursive</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

**Description**

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.
Distinct formulation for emulation of model discrepancies.

**Specification**

**Alias**: paired

**Argument(s)**: none

**Description**

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

**Examples**

Recursive formulation for emulation of model discrepancies.
The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

### Examples

```
7.2.68.1.4.125  expansion_order_sequence

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
```

Sequence of expansion orders used in a multi-stage expansion

### Specification

**Alias:** none  
**Argument(s):** INTEGERLIST  

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Feature</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td><strong>basis_type</strong> Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td><strong>collocation_ratio</strong> Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>expansion_samples_sequence</strong> Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., expansion_samples</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Keywords Area

**import_build_points_file**

| Optional | File containing points you wish to use to build a surrogate |

### Description

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

### See Also

These keywords may also be of interest:

- `expansion_order`

### 7.2.68.1.4.126 dimension_preference

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **dimension_preference**

A set of weights specifying the relative importance of each uncertain variable (dimension)

### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

### Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.1.4.127 basis_type

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Description of Group PCE Basis Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td><strong>Required</strong> (Choose One)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none  
**Argument(s)**: none
Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order \( p \) keep all terms with a \( d \)-dimensional multi index \( \mathbf{i} = (i_1, \ldots, i_d) \) that satisfies
\[
\sum_{k=1}^{d} i_k \leq p
\]

7.2.68.1.4.130 adapted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>advancements</td>
<td></td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td>soft_convergence_limit</td>
<td></td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2 method

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.1.4.133 collocation_ratio

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **collocation_ratio**

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td><strong>orthogonal_matching_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Optional**

<table>
<thead>
<tr>
<th><strong>cross_validation</strong></th>
<th>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ratio_order</strong></td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
</tr>
</tbody>
</table>
### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the `collocation_ratio` allows for specification of a constant factor applied to \( N \) (e.g., `collocation_ratio = 2.` produces samples = \( 2N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using `ratio_order`. In this case, the number of samples becomes \( cN^o \) where \( c \) is the `collocation_ratio` and \( o \) is the `ratio_order`. The `use_derivatives` flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( \frac{cN^o}{n+p+1} \)).

### 7.2.68.1.4.134 collocation_points_sequence

- **Keywords** Area
- **Keywords** method
- **Keywords** bayes_calibration
- **Keywords** queso
- **Keywords** emulator
- **Keywords** ml_pce
- **Keywords** expansion_order_sequence
- **Keywords** collocation_ratio
- **Keywords** collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
Specification

**Alias:** pilot_samples

**Argument(s):** INTEGERLIST

**Description**

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

**See Also**

These keywords may also be of interest:

- `collocation_points`

### 7.2.68.1.4.135 least_squares

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **collocation_ratio**
- **least_squares**

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<td>Description of Group</td>
<td>Dakota Keyword</td>
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Dakota Version 6.15 Reference Manual generated on November 5, 2021
Optional (Choose One)

<table>
<thead>
<tr>
<th>LSQ Regression Approach (Group 1)</th>
<th>svd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>equality_-constrained</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

**Specification**

- **Alias**: none
- **Argument(s)**: none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp  
**Argument(s):** none  
**Child Keywords:**
7.2.68.1.4.139 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias**: bp

**Argument(s)**: none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.1.4.141 basis_pursuit_denoising

- **Keywords Area**
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias**: bpdn

**Argument(s)**: none

**Child Keywords**:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.1.4.142 noise_tolerance

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• basis_pursuit_denoising
• noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.143 least_angle_regression

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.1.4.144 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
7.2.68.1.4.145 least_absolute_shrinkage

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

**Alias**: lasso

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td>noise_tolerance</td>
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<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
<tr>
<td>Optional</td>
<td>l2_penalty</td>
<td>l2_penalty</td>
<td>The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.1.4.146 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**  
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0. (reverts to standard LASSO formulation)

**Description**  
The $l_2$ penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias**: none  
**Argument(s)**: none  

**Child Keywords:**

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<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** ushort max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.1.4.151 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none

Argument(s): REAL

Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
7.2.68.1.4.152 response_scaling

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating **response_scaling**, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- multifidelity_polynomial_chaos
- multifidelity_function_train

7.2.68.1.4.153 use_derivatives

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
Use derivative data to construct surrogate models

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

---

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the import_points file) can be reused.
7.2 method

Specification

Alias: none  
Argument(s): INTEGER  
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.1.4.157 expansion_samples_sequence

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

See Also

These keywords may also be of interest:

- expansion_samples
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.
Specification

**Alias**: import_points_file

**Argument(s)**: STRING

**Default**: no point import from a file

**Child Keywords**: 
The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method
 polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.1.4.160  custom_annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
7.2 method

- import_build_points_file
- custom.annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom.annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

**Resulting tabular file:**

```
%eval_id x1    x2    obj_fn nln_ineq_con_1 nln_ineq_con_2
1   0.9    1.1    0.0002    0.26    0.76
2   0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.14.161 header

• Keywords Area

• method

• bayes_calibration

• queso

• emulator

• ml_pce

• expansion_order_sequence

• import_build_points_file

• custom_annotated

• header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent `custom_annotated`
7.2.68.1.4.162  eval_id

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

7.2.68.1.4.163  interface_id

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.1.4.164 annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- expansion_order_sequence
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom.annotated`
header `eval_id`

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

```
%eval_id interface  x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

### 7.2.68.1.4.165 freeform

- **Keywords Area**
  - method
  - bayes_calibration
  - queso
  - emulator
  - ml_pce
  - expansion_order_sequence
  - import_build_points_file
  - freeform

Selects freeform file format

### Topics

This keyword is related to the topics:

- **file_formats**
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.1.4.166  active_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
• expansion_order_sequence
• import_build_points_file
• active_only

Import only active variables from tabular data file

Topics
This keyword is related to the topics:
  • file_formats

Specification

Alias: none
Argument(s): none

Description
By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.167  orthogonal_least_interpolation

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• ml_pce
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:
### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

7.2.68.1.4.168  `collocation_points_sequence`

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **ml_pce**
- **orthogonal_least_interpolation**
- **collocation_points_sequence**

Sequence of collocation point counts used in a multi-stage expansion

### Specification

**Alias**: none

**Argument(s)**: INTEGERLIST
Description

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`.

See Also

These keywords may also be of interest:

- `collocation_points`

7.2.68.1.4.169 tensor_grid

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
- `emulator`
- `ml_pce`
- `orthogonal_least_interpolation`
- `tensor_grid`

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.1.4.170 reuse_points

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

**import_build_points_file**

- Keywords Area  
- method  
- bayes_calibration  
- queso  
- emulator  
- ml_pce  
- orthogonal_least_interpolation  
- import_build_points_file

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**
Optional (Choose One) | Tabular Format (Group 1) | custom_annotated | Selects custom-annotated tabular file format
---|---|---|---
| | annotated | Selects annotated tabular file format
| | freeform | Selects freeform file format

Optional | active_only | Import only active variables from tabular data file

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```
• **custom.annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

• **file.formats**

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: annotated format

**Child Keywords**: 

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description of Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To control which header row and columns are in the input/output, specify **custom.annotated**, followed by options, in the relevant export/import context.

**Usage Tips**

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to **annotated** format, though **freeform** remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_ id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

### 7.2.68.1.4.173 header

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **ml_pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**
- **custom_annotated**
- **header**

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

**See description of parent** custom_annotated

### 7.2.68.1.4.174 eval_id

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:

```eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  NO_ID  0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.1.4.177 freeform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<p>| | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.68.1.4.178  active_only

- Keywords Area
  - method
  - bayes_calibration
  - queso
  - emulator
  - ml_pce
  - orthogonal_least_interpolation
  - import_build_points_file
  - active_only

Import only active variables from tabular data file

**Topics**

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.179  askey

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- polynomial_chaos  
- askey

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials
Specification

Alias: none
Argument(s): none
Default: PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.1.4.182 export_expansion_file

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

Alias: none
Argument(s): STRING

Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.1.4.183 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- ml_pce
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
7.2 method

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector > 10; else full_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. *diagonal_covariance* is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.1.4.184 full_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mi_pce**
- **full_covariance**

Display the full covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. *full_covariance* is used to force Dakota to output the full covariance matrix.

7.2.68.1.4.185 mf_pce

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**

Multifidelity Polynomial Chaos Expansion as an emulator model.
Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>quadrature_order_sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level_sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>
### Description

Selects a multifidelity polynomial chaos expansion (MF PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF PCE as a surrogate within the Bayesian framework.

### See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no refinement  
**Child Keywords:** none
Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the max_iterations and convergence_tolerance method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

**Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'**.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- p_refinement
- dimension_adaptive

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.1.4.189 sobol

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.
### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** generalized

### Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

### Examples

```plaintext
method,  
    polynomial_chaos  
    sparse_grid_level = 3  
    dimension_adaptive p_refinement sobol  
    max_iterations = 20  
    convergence_tol = 1.e-4
```

### Keywords Area

- method
- bayes_calibration
- queso
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- decay

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

### Specification

**Alias:** none  
**Argument(s):** none

### Description

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.
7.2 method

Examples

    method, polynomial_chaos
        sparse_grid_level = 3
        dimension_adaptive p_refinement decay
        max_iterations = 20
        convergence_tol = 1.e-4

7.2.68.1.4.191 generalized

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none

Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

    method, polynomial_chaos
        sparse_grid_level = 3
        dimension_adaptive p_refinement generalized
        max_iterations = 20
        convergence_tol = 1.e-4

7.2.68.1.4.192 max_refinement_iterations

- Keywords Area
- method
- bayes_calibration
Maximum number of expansion refinement iterations

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.

**Examples**

```plaintext
method,
polynomial_chaos
dimension_adaptive generalized
p_refinement
max_refinement_iterations = 20
convergence_tol = 1.e-4
sparse_grid_level = 1
```

**7.2.68.1.4.193 allocation_control**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **allocation_control**

Sample allocation approach for multifidelity expansions

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**
### Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ an integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

### Default Behavior

The default, when `allocation_control` is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

#### Specification

**Alias**: none

**Argument(s)**: none

### Description

Multifidelity polynomial chaos supports greedy refinement strategies, spanning regression and projection approaches for computing expansion coefficients. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within a unified competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any $z/p/\beta/\beta^*$ level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.
Examples

The following example of greedy multifidelity regression starts from a zeroth-order reference expansion (a constant) for each level, and generates candidate refinements for each level that are competed in an integrated greedy competition. The number of new samples for the incremented candidate expansion order is determined from the collocation ratio. In this case, the number of candidates for each level is limited to one uniform refinement of the current expansion order.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement uniform
    expansion_order_sequence = 0
    collocation_ratio = .9  seed = 160415
    orthogonal_matching_pursuit
    convergence_tolerance 1.e-2
```

The next example employs generalized sparse grids within a greedy multifidelity competition. Each modeling level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all model levels is competed within an integrated greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each model level and the number of candidates grows rapidly with random dimension and grid level.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement dimension_adaptive generalized
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-8
```

7.2.68.1.4.195 discrepancy_emulation

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none

Argument(s): none

Default: distinct

Child Keywords:
7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Discrepancy Emulation Approach (Group 1)</th>
<th>Dakota Keyword</th>
</tr>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>distinct</td>
</tr>
</tbody>
</table>

- **recursive**

<table>
<thead>
<tr>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>distinct</strong></td>
<td>Distinct formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td><strong>recursive</strong></td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

**Description**

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.

**Examples**

```plaintext
method,
  multilevel_polynomial_choas
  expansion_order_sequence = 2
  collocation_ratio = .9
  orthogonal_matching_pursuit
  discrepancy_emulation recursive
```

**7.2.68.1.4.196 distinct**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **discrepancy_emulation**
- **distinct**

Distinct formulation for emulation of model discrepancies.
Specification

Alias: paired
Argument(s): none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.1.4.197 recursive

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples
7.2.68.1.4.198 quadrant_order_sequence

• Keywords Area

• method

• bayes_calibration

• queso

• emulator

• mf_pce

• quadrant_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Each level entry of the quadrant_order_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., quadrant_order

See Also

These keywords may also be of interest:

• quadrant_order
7.2.68.1.4.199  dimension_preference

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- quadrature_order_sequence
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension).

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.1.4.200  nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available
7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Default**: quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are.

7.2.68.1.4.201 non_nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules

Specification

**Alias**: none

**Argument(s)**: none

**Description**

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.1.4.202 sparse_grid_level_sequence

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion
## Specification

**Alias:** none  
**Argument(s):** INTEGERLIST

### Child Keywords:

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<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 1)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>unrestricted</td>
<td>Overide the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

### See Also

These keywords may also be of interest:

- `sparse_grid_level`

7.2.68.1.4.203 `dimension_preference`

- **Keywords Area**
- **method**
A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

**7.2.68.1.4.204 restricted**

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- sparse_grid_level_sequence
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.1.4.205 unrestricted

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• sparse_grid_level_sequence
• unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
7.2 method

7.2.68.1.4.206 nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- sparse_grid_level_sequence
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.207 non_nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- sparse_grid_level_sequence
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none

Argument(s): none
Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

### 7.2.68.1.4.208 expansion_order_sequence

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **expansion_order_sequence**

Sequence of expansion orders used in a multi-stage expansion

#### Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td><strong>expansion_samples_sequence</strong></td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the <code>expansion_samples_sequence</code> applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., <code>expansion_samples</code></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Keywords Area

| Optional | import_build_points_file | File containing points you wish to use to build a surrogate |

#### Description

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

#### See Also

These keywords may also be of interest:

- `expansion_order`

#### 7.2.68.1.4.209 dimension_preference

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- `expansion_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

#### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

#### Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<th>Required/Optional</th>
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<td><strong>Required (Choose One)</strong></td>
<td>PCE Basis Type (Group 1)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>total_order</strong></td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>adapted</strong></td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

**Description**

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none
**Description**

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>advancements</td>
<td>soft_convergence_limit</td>
<td>The maximum number of steps used to expand a basis step. The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.1.4.214 advancements

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.1.4.215 soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.
7.2 method

Specification

Alias: none

Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.1.4.216 collocation_ratio

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none

Argument(s): REAL

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
</tbody>
</table>

Optional (Choose One)

Regression Algorithm (Group 1)
<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>orthogonal_matching_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
</tr>
<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
</tbody>
</table>

**Optional**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>cross_validation</strong></td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
</tr>
<tr>
<td><strong>ratio_order</strong></td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
</tr>
</tbody>
</table>
### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate $N$ from $n$ and $p$, the collocation_ratio allows for specification of a constant factor applied to $N$ (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with $N$ can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $cN^o + 1$).

**7.2.68.1.4.217 collocation_points_sequence**

- **Keywords** Area
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **expansion_order_sequence**
- **collocation_ratio**
- **collocation_points_sequence**

Sequence of collocation point counts used in a multi-stage expansion
Specification

**Alias:** pilot_samples  
**Argument(s):** INTEGERLIST

**Description**

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

**See Also**

These keywords may also be of interest:

- `collocation_points`

7.2.68.1.4.218 least_squares

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
- `emulator`
- `mf_pce`
- `expansion_order_sequence`
- `collocation_ratio`
- `least_squares`

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** svd  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
8.1.4.219 svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp

**Argument(s):** none

**Child Keywords:**
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.1.4.222 noise_tolerance

• Keywords Area
  • bayes_calibration
  • emulator
  • mf_pce
  • expansion_order_sequence
  • collocation_ratio
  • orthogonal_matching_pursuit
  • noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none
Argument(s): REALLIST
Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.223 basis_pursuit

• Keywords Area
  • bayes_calibration
  • emulator
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit \( \ell_1 \)-minimization problem using linear programming.

**Specification**

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit \( \ell_1 \)-minimization problem using linear programming.

7.2.68.1.4.224 basis_pursuit_denoising

- **Keywords Area**

  - method
  - bayes_calibration
  - queso
  - emulator
  - mf_pce
  - expansion_order_sequence
  - collocation_ratio
  - basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising \( \ell_1 \)-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.1.4.225 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.1.4.226 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.1.4.227  noise_tolerance

- Keywords Area

- method

- bayes_calibration

- queso

- emulator

- mf_pce

- expansion_order_sequence

- collocation_ratio

- least_angle_regression

- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
least_absolute_shrinkage

Keywords Area
method
bayes_calibration
quoise
emulator
mf_pce
expansion_order_sequence
colocation_ratio
least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso
Argument(s): none
Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
<tr>
<td>Optional</td>
<td>l2_penalty</td>
<td>l2_penalty</td>
<td></td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
7.2.68.1.4.231 cross_validation

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- cross_validation

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.1.4.232 noise_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.1.4.233 max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** ushort max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.1.4.234 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

**Description**

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use original data  

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

**7.2.68.1.4.236  use_derivatives**

- **Keywords Area**  
- **method**  
- **bayes_calibration**  
- **queso**  
- **emulator**  
- **mf_pce**
7.2 method

- expansion_order_sequence
- collocation_ratio
- use_derivatives

Use derivative data to construct surrogate models.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.1.4.237 tensor_grid

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **expansion_order_sequence**
- **collocation_ratio**
- **tensor_grid**

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.1.4.240 expansion_samples_sequence

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

See Also

These keywords may also be of interest:

- expansion_samples
7.2.68.1.4.241  reuse_points

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• expansion_order_sequence
• expansion_samples_sequence
• reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.68.1.4.242  import_build_points_file

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• expansion_order_sequence
• import_build_points_file

File containing points you wish to use to build a surrogate
7.2 method

Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### Table

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>
• import_build_points_file

• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.68.1.4.244 header

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated
7.2 method

7.2.68.1.4.245 eval_id

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• expansion_order_sequence
• import_build_points_file
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.68.1.4.246 interface_id

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• expansion_order_sequence
• import_build_points_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.1.4.247  annotated

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- expansion_order_sequence
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat' annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn lnl_ineq_con_1 lnl_ineq_con_2
1  NO_ID  0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

Selects freeform file format

Keywords Area

method
bayes_calibration
queso
emulator
mf_pce
expansion_order_sequence
import_build_points_file
freeform

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9   1.1   0.0002   0.26   0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.1.4.249 active_only

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
• expansion_order_sequence
• import_build_points_file
• active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.250 orthogonal_least_interpolation

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:
### Keywords Area

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td><code>collocation_points_sequence</code></td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>tensor_grid</code></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>reuse_points</code></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>import_build_points_file</code></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

#### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data

7.2.68.1.4.251 `collocation_points_sequence`

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- orthogonal_least_interpolation
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

#### Specification

**Alias:** none

**Argument(s):** INTEGERLIST
Description

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

See Also

These keywords may also be of interest:

- `collocation_points`

7.2.68.1.4.252 tensor_grid

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

### Specification

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

### Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```bash
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

---

<table>
<thead>
<tr>
<th>Optional</th>
<th>Tabular Format (Group 1)</th>
<th>custom_annotated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
- **custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```python
evironment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

### 7.2.68.1.4.256 header

- **Keywords Area**
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent custom_annotated

### 7.2.68.1.4.257 eval_id

- **Keywords Area**
- method
- bayes_calibration
- queso
Enable evaluation ID column in custom-annotated tabular file

**Specification**

*Alias:* none  
*Argument(s):* none  
*Default:* no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

*Alias:* none  
*Argument(s):* none  
*Default:* no interface_id column

**Description**

See description of parent `custom_annotated`
7.2.68.1.4.259 annotated

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_pce
• orthogonal_least_interpolation
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.1.4.260  freeform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

**7.2.68.1.4.261 active_only**

- Keywords Area

- method

- bayes_calibration

- queso

- emulator

- mf_pce

- orthogonal_least_interpolation

- import_build_points_file

- active_only

Import only active variables from tabular data file

**Topics**

This keyword is related to the topics:

- file_formats
By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.1.4.262 askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

These keywords may also be of interest:

- polynomial_chaos
- wiener
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)

### Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

### See Also

These keywords may also be of interest:

- `polynomial_chaos`
- `askey`

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
Specification

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.1.4.265 export_expansion_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **export_expansion_file**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

**Alias:** none

**Argument(s):** STRING

Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.1.4.266 diagonal_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **diagonal_covariance**

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector > 10; else full_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.1.4.267  full_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_pce**
- **full_covariance**

Display the full covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.

7.2.68.1.4.268  sc

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **sc**

Stochastic Collocation as an emulator model.
Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**: 
### Required/Optional

**Optional (Choose One)**

<table>
<thead>
<tr>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Employ h-refinement to refine the grid</td>
<td>h_refinement</td>
<td></td>
</tr>
<tr>
<td>Maximum number of expansion refinement iterations</td>
<td>max_refinement_-iterations</td>
<td></td>
</tr>
<tr>
<td>Order for tensor-products of Gaussian quadrature rules</td>
<td>quadrature_order</td>
<td></td>
</tr>
<tr>
<td>Level to use in sparse grid integration or interpolation</td>
<td>sparse_grid_level</td>
<td></td>
</tr>
<tr>
<td>Use piecewise local basis functions</td>
<td>piecewise</td>
<td></td>
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</table>

**Required (Choose One)**

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<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Interpolation Grid Type (Group 2)</td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
</tbody>
</table>

**Optional (Choose One)**

<table>
<thead>
<tr>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basis Polynomial Family (Group 3)</td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>----------</td>
<td>-----------------</td>
<td>---------------------------------------------------</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 4)</td>
<td>diagonal_covariance</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

**Description**

Selects stochastic collocation (SC) model to use in the Bayesian likelihood calculations. Most specification options are carried over for using SC as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- `stoch_collocation`

**Automatic polynomial order refinement**

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no refinement

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group p-refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>uniform</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
</tbody>
</table>
Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

### Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidate sets, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

### 7.2.68.1.4.270 uniform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.
Specification

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.1.4.271 dimension_adaptive

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **sc**
- **p_refinement**
- **dimension_adaptive**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Dimension Adaptivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
<td></td>
</tr>
</tbody>
</table>
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

### 7.2.68.1.4.272 sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **sc**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

**Specification**

- **Alias:** none
- **Argument(s):** none
- **Default:** generalized

**Description**

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```

---

**7.2.68.1.4.273  h_refinement**

- **Keywords Area**
- **method**
- **bayes_calibration**

---

**7.2.68.1.4.274  h_refinement**

- **Keywords Area**
- **method**
- **bayes_calibration**
• queso
• emulator
• sc
• h_refinement

Employ h-refinement to refine the grid

Specification

Alias: none
Argument(s): none
Default: no refinement

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>h-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

Description

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_adaptive options.
Refine an expansion uniformly in all dimensions.

**Specification**

*Alias:* none

*Argument(s):* none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

**Performance**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

*Alias:* none

*Argument(s):* none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
### Required (Choose One)

**Dimension Adaptivity Estimation Approach (Group 1)**

- **sobol**
  - Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

- **generalized**
  - Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

#### 7.2.68.1.4.277 sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **sc**
- **h_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.1.4.278 generalized

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- h_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none

Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

Examples

```plaintext
method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
Planned future capability for local pointwise refinement within a generalized sparse grid.

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

**Maximum number of expansion refinement iterations**

**Specification**

**Alias**: none  
**Argument(s)**: INTEGER  
**Default**: 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

```
method,
polynomial_chaos
dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

7.2.68.1.4.281 quadrature_order

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

Specification

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_-preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order, and, optionally, dimension_preference). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a dimension_preference,
where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified `quadrature_order` and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n}(p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the `quadrature_order` specification supports an array input.

A corresponding sequence specification is documented at, e.g., `quadrature_order_sequence` and `quadrature_order_sequence`

See Also

These keywords may also be of interest:

- `quadrature_order_sequence` and `method-multifidelity_stoch_collocation-quadrature_order_sequence`

7.2.68.1.4.282 dimension_preference

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- quadrature_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.1.4.283  nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- quadrature_order
- nested

Enforce use of nested quadrature rules if available

Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.284  non_nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules
7.2 method

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.1.4.285 sparse_grid_level

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tr>
<td>Optional</td>
<td>dimension_preference</td>
<td>nodal</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension) Employ a nodal sparse grid construction in stochastic collocation</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Group 1</td>
<td>nodal</td>
<td></td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with sparse_grid_level and, optionally, dimension_preference). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the dimension_preference specification enables the use of anisotropic sparse grids (refer to the PCE description in the User’s Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full sparse_grid_level and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User’s Manual for additional details). As for quadrature_order, the sparse_grid_level specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., sparse_grid_level_sequence and sparse_grid_level_sequence.

See Also

These keywords may also be of interest:

- sparse_grid_level_sequence method-multifidelity_stoch_collocation-sparse_grid_level_sequence

7.2.68.1.4.286 dimension_preference

- Keywords Area
- method
A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

**Employ a nodal sparse grid construction in stochastic collocation**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** nodal
Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.

7.2.68.1.4.288  hierarchical

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- sparse_grid_level
- hierarchical

Employ a hierarchical sparse grid construction

Specification

Alias: none
Argument(s): none

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.68.1.4.289  restricted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.
7.2.68.1.4.290  unrestricted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive_p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences. To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.1.4.291 nested

* Keywords Area
* method
* bayes_calibration
* quiero
* emulator
* sc
* sparse_grid_level
* nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.292 non_nested

* Keywords Area
* method
* bayes_calibration
* quiero
* emulator
* sc
* sparse_grid_level
* non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none
### Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.681.4.293 piecewise

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- piecewise

Use piecewise local basis functions

### Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

### Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.1.4.294 askey

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

7.2.68.1.4.295 wiener

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
7.2 method

See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.1.4.296 use_derivatives

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.1.4.297 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- sc
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
Specification

Alias: none
Argument(s): none
Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. \textit{diagonal\_covariance} is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.1.4.298  full\_covariance

• Keywords Area
• method
• bayes\_calibration
• qeso
• emulator
• sc
• full\_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. \textit{full\_covariance} is used to force Dakota to output the full covariance matrix.

7.2.68.1.4.299  mf\_sc

• Keywords Area
• method
• bayes\_calibration
• qeso
• emulator
• mf\_sc

Multifidelity Stochastic Collocation as an emulator model.
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td><strong>Optional (Choose One)</strong></td>
<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
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<tr>
<td>Optional</td>
<td></td>
<td>h_refinement</td>
<td>Employ h-refinement to refine the grid</td>
</tr>
<tr>
<td>Optional</td>
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<td>max_refinement_-iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
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<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_- emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
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<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Interpolation Grid Type (Group 2)</td>
<td>quadrature_order_-sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
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<td></td>
<td>sparse_grid_level_-sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td><strong>Optional (Choose One)</strong></td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
</tbody>
</table>
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Optional**

<table>
<thead>
<tr>
<th>Optional/Choose One</th>
<th>Covariance Type (Group 4)</th>
<th>Use derivaives</th>
<th>Use derivative data to construct surrogate models</th>
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<tr>
<td></td>
<td>diagonal,covariance</td>
<td></td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td></td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

**Description**

Selects a multifidelity stochastic collocation (MF SC) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF SC as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- **multifidelity_stoch_collocation**

7.2.68.1.4.300  p_refinement

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- p_refinement

Automatic polynomial order refinement

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Dakota Keyword Description

<table>
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<tr>
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<th>Description of Group p-refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>uniform</td>
<td>dimension_adaptive</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
</tbody>
</table>

### Description

The **p_refinement** keyword specifies the usage of automated polynomial order refinement, which can be either **uniform** or **dimension_adaptive**.

The **dimension_adaptive** option is supported for the tensor-product quadrature and Smolyak sparse grid options and **uniform** is supported for tensor and sparse grids as well as regression approaches (**collocation_points** or **collocation_ratio**).

Each of these refinement cases makes use of the **max_iterations** and **convergence_tolerance** method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The **dimension_adaptive** case can be further specified to utilize **sobol**, **decay**, or **generalized** refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (**sobol** case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (**decay** case: low decay rates result in high dimension preference). In these two cases as well as the **uniform** refinement case, the **quadrature_order** or **sparse_grid_level** are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the **uniform** refinement case with regression approaches, the **expansion_order** is ramped by one on each iteration while the oversampling ratio (either defined by **collocation_ratio** or inferred from **collocation_points** based on the initial expansion) is held fixed. Finally, the generalized **dimension_adaptive** case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of **p_refinement** or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

**7.2.68.1.4.301 uniform**

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
Refine an expansion uniformly in all dimensions.

Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.
<table>
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<tr>
<th>Required/Optional</th>
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</tr>
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<tr>
<td>Required (Choose One)</td>
<td></td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.1.4.303 sobol

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** generalized
7.2 method

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.1.4.304 generalized

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_sc
• p_refinement
• dimension_adaptive
• generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none
Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
7.2.68.1.4.305 \( h_{\text{refinement}} \)

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- \( h_{\text{refinement}} \)

Employ \( h_{\text{refinement}} \) to refine the grid.

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no refinement

**Child Keywords**:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<tr>
<td><strong>Dakota Keyword</strong></td>
<td><strong>Dakota Keyword Description</strong></td>
</tr>
<tr>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td>dimension_adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

**Description**

Automated expansion refinement can be selected as either \( p_{\text{refinement}} \) or \( h_{\text{refinement}} \), and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The \( h_{\text{refinement}} \) specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of \( h_{\text{refinement}} \)). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_adaptive options.
7.2.68.1.4.306 uniform

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- h_refinement
- uniform

Refine an expansion uniformly in all dimensions.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.1.4.307 dimension_adaptive

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- h_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.1.4.308 sobol

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- h_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

### Specification

- **Alias:** none
- **Argument(s):** none
- **Default:** generalized
7.2 method

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.1.4.309 generalized

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- h_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none
Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
Planned future capability for local pointwise refinement within a generalized sparse grid.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
7.2 method

Examples

method,
  polynomial_chaos
    dimension_adaptive generalized
    p_refinement
      max_refinement_iterations = 20
      convergence_tol = 1.e-4
      sparse_grid_level = 1

7.2.68.1.4.312 allocation_control

  • Keywords Area
  • method
  • bayes_calibration
  • queso
  • emulator
  • mf_sc
  • allocation_control

Sample allocation approach for multifidelity expansions

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Multifidelity Sample Allocation Control (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Required</td>
<td>Description</td>
<td>greedy</td>
<td>Sample allocation based on greedy refinement within multifidelity stochastic collocation</td>
</tr>
</tbody>
</table>

Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

Default Behavior

The default, when allocation_control is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).
Sample allocation based on greedy refinement within multifidelity stochastic collocation

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Multifidelity stochastic collocation supports greedy refinement strategies using tensor and sparse grids for both nodal and hierarchical collocation approaches. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within an integrated competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.

**Examples**

The following example of greedy multifidelity stochastic collocation using nodal interpolation starts from a zeroth-order expansion (a constant) for each level, and generates uniform candidate refinements for each level that are competed in a greedy competition. The number of new samples for the incremented candidate expansion order is determined from the quadrature rules of the new sparse grid level. In this case, the number of candidates for each level is limited to one uniform refinement of the current sparse grid level.

```plaintext
method,
   model_pointer = 'HIERARCH'
   multifidelity_stoch_collocation
      nodal
      allocation_control greedy
      p_refinement uniform
      sparse_grid_level_sequence = 0 unrestricted
      convergence_tolerance 1.e-3
```

The next example employs generalized sparse grids and hierarchical interpolation. Each level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all levels is competed within a unified greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each level and the number of candidates grows rapidly with random dimension and grid level.
7.2 method

method,
model_pointer = 'HIERARCH'
multifidelity_stoch_collocation
  hierarchical
  allocation_control greedy
  p_refinement dimension Adaptive generalized
  sparse_grid_level_sequence = 0 unrestricted
  convergence_tolerance 1.e-8

7.2.68.1.4.314 discrepancy_emulation

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_sc
• discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none
Default: distinct
Child Keywords:

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Discrepancy Emulation Approach (Group 1)</th>
<th>Dakota Keyword distinct</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td>Distinct formulation for emulation of model discrepancies.</td>
<td>recursive</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. distinct emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. recursive emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.
The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

Examples

```
method,
    multilevel_polynomial_chaos
    expansion_order_sequence = 2
    collocation_ratio = .9
    orthogonal_matching_pursuit
    discrepancy_emulation recursive
```

7.2.68.1.4.315  distinct

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- discrepancy_emulation
- distinct

Distinct formulation for emulation of model discrepancies.

Specification

**Alias:** paired

**Argument(s):** none

Description

The **distinct** approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.1.4.316  recursive

- Keywords Area
- method
- bayes_calibration
Recursive formulation for emulation of model discrepancies.

Specification

Alias: none

Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.1.4.317 quadrature_order_sequence

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_sc
• quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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Dakota Version 6.15 Reference Manual generated on November 5, 2021
<table>
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<tr>
<th>Optional</th>
<th>Quadrature Rule Nesting (Group 1)</th>
<th>dimension_preference</th>
<th>A set of weights specifying the relative importance of each uncertain variable (dimension)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Choose One</td>
<td>nested</td>
<td>non_nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
</tbody>
</table>

**Description**

Each level entry of the `quadrature_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `quadrature_order`

**See Also**

These keywords may also be of interest:

- `quadrature_order`

7.2.68.1.4.318  `dimension_preference`

- `Keywords Area`
- `method`
- `bayes_calibration`
- `queso`
- `emulator`
- `mf_sc`
- `quadrature_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids
Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.1.4.319 nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.1.4.320 non_nested

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules
Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.1.4.321  sparse_grid_level_sequence

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
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<td></td>
<td>dimension_ - preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Group 1</td>
<td>nodal</td>
<td>Employ a nodal sparse grid construction in stochastic collocation</td>
</tr>
</tbody>
</table>
**Optional (Choose One)** | **Quadrature Rule Growth (Group 2)** | **restricted** | Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

**Optional (Choose One)** | **Quadrature Rule Nesting (Group 3)** | **nested** | Enforce use of nested quadrature rules if available

**Optional (Choose One)** | **Quadrature Rule Nesting (Group 3)** | **non_nested** | Enforce use of non-nested quadrature rules

---

### Description

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

### See Also

These keywords may also be of interest:

- `sparse_grid_level`

---

7.2.68.1.4.322  **dimension_preference**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_sc**
- `sparse_grid_level_sequence`
- **dimension_preference**

A set of weights specifying the relative importance of each uncertain variable (dimension)
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- **sobol**
- **decay**

7.2.68.1.4.323 nodal

- **Keywords Area**
- **method**
- **bayes_calibration**
- **queso**
- **emulator**
- **mf_sc**
- **sparse_grid_level_sequence**
- **nodal**

Employ a nodal sparse grid construction in stochastic collocation

Specification

**Alias:** none

**Argument(s):** none

**Default:** nodal

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.
7.2.68.1.4.324  hierarchical

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_sc
• sparse_grid_level_sequence
• hierarchical

Employ a hierarchical sparse grid construction

Specification

Alias: none
Argument(s): none

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.68.1.4.325  restricted

• Keywords Area
• method
• bayes_calibration
• queso
• emulator
• mf_sc
• sparse_grid_level_sequence
• restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

Alias: none
Argument(s): none
Default: restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.1.4.326 unrestricted

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none
Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none
Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.1.4.329  piecewise

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- piecewise

Use piecewise local basis functions

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.1.4.330  askey

- Keywords Area
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.1.4.332 use_derivatives

Use derivative data to construct surrogate models

Specification

**Alias:** none

**Argument(s):** none

**Default:** use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.1.4.333 diagonal_covariance

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

**Aliases**: none

**Argument(s)**: none

**Default**: diagonal_covariance for response vector > 10; else full_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.1.4.334  full_covariance

- **Keywords Area**
- method
- bayes_calibration
- queso
- emulator
- mf_sc
- full_covariance

Display the full covariance matrix

7.2.68.1.5  standardized_space

- **Keywords Area**
- method
- bayes_calibration
- queso
- standardized_space

Perform Bayesian inference in standardized probability space

**Specification**

**Aliases**: none

**Argument(s)**: none

**Description**

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.
Description

This option transforms the inference process (MCMC sampling and any emulator model management) into a standardized probability space.

The variable transformations performed are as described in askey.

Default Behavior

The default for the Gaussian process and no emulator options is to perform inference in the original probability space (no transformation). Polynomial chaos and stochastic collocation emulators, on the other hand, are always formed in standardized probability space, such that the inference process is also performed in this standardized space.

Expected Output

The user will see the truth model evaluations performed in the original space, whereas any method diagnostics relating to the MCMC samples (e.g., QUESO data in the outputData directory) will report points and response data (response gradients and Hessians, if present, will differ but response values will not) that correspond to the transformed space.

Usage Tips

Selecting standardized_space generally has the effect of scaling the random variables to be of comparable magnitude, which can improve the efficiency of the Bayesian inference process.

Examples

```
method,
  bayes_calibration queso
  samples = 2000 seed = 348
dram
  standardized_space
```

7.2.68.1.6 logit_transform

- Keywords Area
- method
- bayes_calibration
- queso
- logit_transform

Utilize the logit transformation to reduce sample rejection for bounded domains

Specification

Alias: none

Argument(s): none

Description

The logit transformation performs an internal variable transformation from bounded domains to unbounded domains in order to reduce sample rejection due to an out-of-bounds condition.

Default Behavior

This option is experimental at present, and is therefore defaulted off.
Usage Tips

This option can be helpful when regions of high posterior density exist in the corners of a multi-dimensional bounded domain. In these cases, it may be difficult to generate feasible samples from the proposal density, such that transformation to unbounded domains may greatly reduce sample rejection rates.

Examples

```yaml
method,
bayes_calibration queso
samples = 2000 seed = 348
dram
logit_transform
```

7.2.68.1.7 export_chain_points_file

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file

Export the MCMC chain to the specified filename

Specification

Alias: none

Argument(s): STRING

Default: chain export to default filename

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

The filename to which the final MCMC posterior chain will be exported.

Default Behavior  No export to file.

Expected Output

A tabular data file will be produced in the specified format (annotated by default) containing samples from the posterior distribution.

Usage Tips
Additional Discussion

7.2.68.1.7.1 custom_annotated

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior
The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id    x1    x2    obj_fn    nln_ineq_con_1    nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.1.7.2 header

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none  
**Argument(s):** none

Description

See description of parent custom_annotated
7.2.68.1.7.3 eval_id

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.1.7.4 interface_id

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated
7.2.68.1.7.5 annotated

- **Keywords Area**
  - method
  - bayes_calibration
  - queso
  - export_chain_points_file
  - annotated

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
... 7.2.68.1.7.6 freeform
```

- Keywords Area
- method
- bayes_calibration
- queso
- export_chain_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips
• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
    tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
  ...
```

use the DRAM MCMC algorithm

Topics

This keyword is related to the topics:

• bayesian_calibration

Specification

**Alias:** none

**Argument(s):** none

**Default:** dram
Description

The type of Markov Chain Monte Carlo used. This keyword specifies the use of DRAM, (Delayed Rejection Adaptive Metropolis)[40].

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Usage Tips

If the user knows very little about the proposal covariance, using dram is a recommended strategy. The proposal covariance is adaptively updated, and the delayed rejection may help improve low acceptance rates.

Examples

```
method, bayes_calibration queso
dram
samples = 10000 seed = 348
```

7.2.68.1.9 delayed_rejection

- Keywords Area
- method
- bayes_calibration
- queso
- delayed_rejection

Use the Delayed Rejection MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

**Alias:** none

**Argument(s):** none

**Default:** dram

Description

This keyword specifies the use of the Delayed Rejection algorithm in which there can be a delay in rejecting samples from the chain. That is, the "DR" part of DRAM is used but the "AM" part is not, rather a regular Metropolis-Hastings algorithm is used.

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.
7.2 method

Usage Tips

If the user knows something about the proposal covariance or the proposal covariance is informed through derivative information, using delayed_rejection is preferred over dram: the proposal covariance is already being informed by derivative information and the adaptive metropolis is not necessary.

Examples

```
method,
   bayes_calibration queso
delayed_rejection
samples = 10000 seed = 348
```

See Also

These keywords may also be of interest:

- proposal_covariance

7.2.68.1.10 adaptive_metropolis

- Keywords Area
- method
- bayes_calibration
- queso
- adaptive_metropolis

Use the Adaptive Metropolis MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

**Alias:** none

**Argument(s):** none

**Default:** dram

Description

This keyword specifies the use of the Adaptive Metropolis algorithm. That is, the "AM" part of DRAM is used but the "DR" part is not: specifying this keyword activates only the Adaptive Metropolis part of the MCMC algorithm, in which the covariance of the proposal density is updated adaptively.

Default Behavior

Five MCMC algorithm variants are supported in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.
Four MCMC variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings, and adaptive_metropolis.

Usage Tips
If the user knows very little about the proposal covariance, but doesn’t want to incur the cost of using full dram with both delayed rejection and adaptive metropolis, specifying only adaptive_metropolis offers a good strategy.

Examples
```
    method,  bayes_calibration queso
                 adaptive_metropolis
                   samples = 10000 seed = 348
```

7.2.68.1.11 metropolis_hastings

- Keywords Area
- method
- bayes_calibration
- queso
- metropolis_hastings

Use the Metropolis-Hastings MCMC algorithm

Topics
This keyword is related to the topics:
- bayesian_calibration

Specification

Alias: none
Argument(s): none
Default: dram

Description
This keyword specifies the use of a Metropolis-Hastings algorithm for the MCMC chain generation. This means there is no delayed rejection and no adaptive proposal covariance updating as in DRAM.

Default Behavior
Five MCMC algorithm variants are supported currently in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Four MCMC algorithm variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings, and adaptive_metropolis.

Usage Tips
If the user wants to use Metropolis-Hastings, possibly as a comparison to the other methods which involve more chain adaptation, this is the MCMC type to use.
Examples

```
method, bayes_calibration queso
  metropolis_hastings
  samples = 10000 seed = 348
```

7.2.6.8.1.12 multilevel

- Keywords Area
- method
- bayes_calibration
- queso
- multilevel

Use the multilevel MCMC algorithm.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

**Alias:** none

**Argument(s):** none

**Default:** dram

Description

Selects the multilevel algorithm described in[73].

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Usage Tips

The multilevel algorithm is a more experimental algorithm than the other MCMC approaches mentioned above. It works well in cases where the prior can be “evolved” to a posterior in a structured way. Currently, the multilevel option is not in production form.

Examples

```
method, bayes_calibration queso
  multilevel
  samples = 10000 seed = 348
```
7.2.68.1.13 pre_solve

- Keywords Area
- method
- bayes_calibration
- queso
- pre_solve

Perform deterministic optimization for MAP before Bayesian calibration

Specification

**Alias:** none

**Argument(s):** none

**Default:** nip pre-solve for emulators

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Pre-solve Optimizer (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required One</strong></td>
<td></td>
<td>sqp</td>
<td>Uses a sequential quadratic programming method for underlying optimization</td>
</tr>
<tr>
<td>nip</td>
<td></td>
<td></td>
<td>Uses a nonlinear interior point method for underlying optimization</td>
</tr>
<tr>
<td>none</td>
<td></td>
<td></td>
<td>Deactivates MAP pre-solve prior to initiating the MCMC process</td>
</tr>
</tbody>
</table>

**Description**

When specified, Dakota will perform a deterministic derivative-based optimization to maximize the log posterior (minimize the negative log posterior = misfit - log_prior + constant normalization factors). The Markov chain in Bayesian calibration will subsequently be started at the best point found in the optimization (the MAP point), which can eliminate the need for "burn in" of the chain in which some initial portion of the chain is discarded. Note that both optimization methods available (sqp and nip) require derivatives of the negative log posterior, either first-order in the case of SQP (with quasi-Newton Hessians from secant updates) or second-order in the case of full-Newton NIP (with explicit Hessian use). The derivatives will be computed from the same model used for the MCMC process; e.g. if an emulator is used, the emulator derivatives will be used, otherwise they will be based on the user's model specification for the model.

It is important to clarify that the use of the Hessian of the negative log posterior within a full Newton solver does not strictly require Hessians from the model response quantities of interest (QoIs). Rather, the Hessian of the negative log posterior is formed from an exact Hessian of the negative log prior and a misfit Hessian that can be either exact or approximated: the full misfit Hessian can be formed using QoI residuals, gradients, and Hessians or the
Gauss-Newton approximate misfit Hessian can be formed using only QoI gradients [16]. This Hessian composition is configured automatically based on MAP solver selection and the emulator’s or simulation model’s support for derivatives.

**Default Behavior** The default MAP pre-solve behavior depends on the use of an emulator model within the inference process.

If there is an emulator (for which derivatives are easily computed), then the MAP pre-solve is active by default and a full Newton NIP formulation is selected if OPT++ is available. The default use of a MAP pre-solve can be overridden using “pre_solve none” and the default selection of OPT++ full Newton NIP can be replaced with NPSOL SQP using “pre_solve sqp.” Depending on the emulator’s support for derivatives of the simulated QoI (gradients for dakota GP and stochastic collocation; gradients and Hessians for surfpack GP and polynomial chaos), the contribution of the misfit Hessian to the Hessian of the negative log posterior will be computed either using the full misfit Hessian or its Gauss-Newton approximation (refer to Bayesian chapter in [16]).

If no emulator model is specified, then the pre-solve is bypassed by default and the MCMC chain is initiated from user-specified (or default) initial value for the prior distributions of the random variables. This default can be overridden by specifying “pre_solve nip” for a full Newton NIP solution or “pre_solve sqp” for an NPSOL SQP solution. Both MAP pre-solve approaches require QoI gradients from the simulation model, and the full Newton approach can further leverage QoI Hessians when available (though not required due to the Gauss-Newton approximation, as explained previously).

**Expected Output** When pre-solve is enabled, the output will include a deterministic optimization, followed by a Bayesian calibration. The final results will include the MAP point as well as posterior statistics from the MCMC chain. The MAP point that is reported is the point with highest posterior probability spanning both the pre-solve and the subsequent MCMC chain; it will most commonly reflect the end state of the pre-solve, although it can reflect subsequent improvements from the chain evolution, should they occur.

**Examples**

```r
method
  bayes_calibration queso
  samples = 2000 seed = 348
  delayed_rejection
  emulator
    pce sparse_grid_level = 2
    pre_solve nip # default for emulators
```

7.2.68.1.13.1 sqp

- Keywords Area
- method
- bayes_calibration
- queso
- pre_solve
- sqp

Uses a sequential quadratic programming method for underlying optimization

**Specification**

**Alias**: none

**Argument(s)**: none
Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The sqp keyword directs Dakota to use a sequential quadratic programming method to solve that problem. A sequential quadratic programming solves a sequence of linearly constrained quadratic optimization problems to arrive at the solution to the optimization problem.

7.2.68.1.13.2 nip

- Keywords Area
- method
- bayes_calibration
- queso
- pre_solve
- nip

Uses a nonlinear interior point method for underlying optimization

Specification

Alias: none
Argument(s): none

Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The nip keyword directs Dakota to use a nonlinear interior point to solve that problem. A nonlinear interior point method traverses the interior of the feasible region to arrive at the solution to the optimization problem.

7.2.68.1.13.3 none

- Keywords Area
- method
- bayes_calibration
- queso
- pre_solve
- none

Deactivates MAP pre-solve prior to initiating the MCMC process.

Specification

Alias: none
Argument(s): none
Description

Pre-solving for the maximum a posteriori probability (MAP) point could be undesirable when testing MCMC performance or too expensive to pursue in some settings. The option "none" provides an override that deactivates the pre-solve option for cases where it would normally be active by default (e.g., for emulator models).

7.2.68.1.14 proposal_covariance

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance

Defines the technique used to generate the MCMC proposal covariance.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Proposal Covariance Source (Group 1)</td>
<td>prior</td>
<td>Uses the covariance of the prior distributions to define the MCMC proposal covariance.</td>
</tr>
<tr>
<td>derivatives</td>
<td></td>
<td></td>
<td>Use derivatives to inform the MCMC proposal covariance.</td>
</tr>
<tr>
<td>values</td>
<td></td>
<td></td>
<td>Specifies matrix values to use as the MCMC proposal covariance.</td>
</tr>
</tbody>
</table>
**Description**

The proposal covariance is used to define a multivariate normal (MVN) jumping distribution used to create new points within a Markov chain. That is, a new point in the chain is determined by sampling within a MVN probability density with prescribed covariance that is centered at the current chain point. The accuracy of the proposal covariance has a significant effect on rejection rates and the efficiency of chain mixing.

**Default Behavior**

The default proposal covariance is `prior` when no emulator is present; `derivatives` when an emulator is present.

**Expected Output**

The effect of the proposal covariance is reflected in the MCMC chain values and the rejection rates, which can be seen in the diagnostic outputs from the QUESO solver within the `QuesoDiagnostics` directory.

**Usage Tips**

When derivative information is available inexpensively (e.g., from an emulator model), the derived-based proposal covariance forms a more accurate proposal distribution, resulting in lower rejection rates and faster chain mixing.

7.2.68.1.14.1 `prior`

- **Keywords Area**
- `method`
- `bayes_calibration`
- `queso`
- `proposal_covariance`
- `prior`

Uses the covariance of the prior distributions to define the MCMC proposal covariance.

**Topics**

This keyword is related to the topics:

- `bayesian_calibration`

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:** none
### Description

This keyword selection results in definition of the MCMC proposal covariance from the covariance of the prior distributions. This covariance is currently assumed to be diagonal without correlation.

### Default Behavior

This is the default proposal_covariance option.

### Usage Tips

Since this proposal covariance is defined globally, the chain does not need to be periodically restarted using local updates to this proposal. However, it is usually effective to adapt the proposal using one of the adaptive metropolis MCMC options.

### Examples

```plaintext
method,
bayes_calibration queso
  samples = 2000 seed = 348
dram
  proposal_covariance prior
```

### Specification

- **Alias:** none
- **Argument(s):** REAL
- **Default:** 1.0

### Description

The initial proposal covariance will be given by the prior variance times the multiplier.

**Default Behavior** When using prior-based proposal covariance, the default is to use the prior variance for the proposal (multiplier = 1.0)
Usage Tips

The prior variance may result in too tight or narrow a proposal covariance. The multiplier can be used to scale all entries of the prior variance in determining the initial proposal covariance.

Examples

method
  bayes_calibration queso
  samples = 2000 seed = 348
  dram
  proposal_covariance prior
  multiplier 0.1

7.2.68.1.14.3 derivatives

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- derivatives

Use derivatives to inform the MCMC proposal covariance.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>update_period</td>
<td>Period at which to update derivative-based proposal covariance</td>
</tr>
</tbody>
</table>

Description

This keyword selection results in definition of the MCMC proposal covariance from the Hessian of the misfit function (negative log likelihood), where this Hessian is defined from either a Gauss-Newton approximation (using only first derivatives of the calibration terms) or a full Hessian (using values, first derivatives, and second derivatives of the calibration terms). If this Hessian is indeterminate, it will be corrected as described in[16].
Default Behavior  The default is prior based proposal covariance. This is a more advanced option that exploits structure in the form of the likelihood.

Expected Output

When derivatives are specified for defining the proposal covariance, the misfit Hessian and its inverse (the MVN proposal covariance) will be output to the standard output stream.

Usage Tips

The full Hessian of the misfit is used when either supported by the emulator in use (for PCE and surfpack GP, but not SC or dakota GP) or by the user’s response specification (Hessian type is not "no_hessians"), in the case of no emulator. When this full Hessian is indefinite and cannot be inverted to form the proposal covariance, fallback to the positive semi-definite Gauss-Newton Hessian is employed.

Since this proposal covariance is locally accurate, it should be updated periodically using the update_period option. While the adaptive metropolis option can be used in combination with derivative-based preconditioning, it is generally preferable to instead decrease the proposal update period due to the improved local accuracy of this approach.

Examples

Generate a 2000 sample posterior chain, using derivatives to initialize the proposal covariance at the start of the chain.

```
method, bayes_calibration queso
    samples = 2000 seed = 348
    delayed_rejection
    emulator pce sparse_grid_level = 2
    proposal_covariance derivatives
```

7.2.68.1.14.4 update_period

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- derivatives
- update_period

Period at which to update derivative-based proposal covariance

Specification

Alias: none

Argument(s): INTEGER

Description

For derivative-based proposal covariance, this specifies the period (number of accepted MCMC samples) after which the proposal covariance is updated using derivative values at the current chain point.
When `update_period` is not specified, derivatives will inform the proposal covariance at the start of the chain, but not updated further.

**Usage Tips**

The `update_period` should be tailored to the size of the total chain, accounting for the relative expense of derivative-based proposal updates.

**Examples**

```
method, bayes_calibration queso
   samples = 2000 seed = 348
   delayed_rejection
   emulator pce sparse_grid_level = 2
   proposal_covariance derivatives
      update_period = 40  # update proposal covariance every 40 points
```

7.2.68.1.14.5 values

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- values

Specifies matrix values to use as the MCMC proposal covariance.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Values For (Group 1)</th>
<th>Dakota Keyword diagonal</th>
</tr>
</thead>
</table>
| Specify the diagonal matrix format when specifying a user-specified proposal covariance.
7.2 method

| matrix | Specifies the full matrix format when specifying a user-specified proposal covariance. |

Description

This keyword selection results in definition of the MCMC proposal covariance from user-specified matrix values. The matrix input format must be declared as either a full matrix or a matrix diagonal.

Default Behavior

This option is not the default, and generally implies special a priori knowledge from the user.

Usage Tips

This option is not supported for the case of transformations to standardized probability space.

Examples

```
method, bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
    values ... # See leaf nodes for required format option
```

7.2.68.1.14.6 diagonal

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- values
- diagonal

Specifies the diagonal matrix format when specifying a user-specified proposal covariance.

Specification

Alias: none

Argument(s): none

Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.
Examples

```
method,
  bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
    diagonal values 1.0e6 1.0e-1
```

7.2.68.1.14.7 matrix

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- values
- matrix

Specifies the full matrix format when specifying a user-specified proposal covariance.

Specification

**Alias:** none

**Argument(s):** none

Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.

Examples

```
method,
  bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
    matrix values 1.0 0.1
                0.1 2.0
```

7.2.68.1.14.8 filename

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- filename

Uses a file to import a user-specified MCMC proposal covariance.
Specification

**Alias:** none

**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Values For (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>diagonal</td>
<td>Specifies the diagonal matrix format when importing a user-specified proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>matrix</td>
<td>Specifies the full matrix format when importing a user-specified proposal covariance.</td>
</tr>
</tbody>
</table>

**Description**

This keyword selection results in definition of the MCMC proposal covariance from importing data a user-specified filename. This import must be declared as either a full matrix or a matrix diagonal.

**Default Behavior**

This option is not the default, and generally implies special a priori knowledge from the user.

**Usage Tips**

This option is not supported for the case of transformations to standardized probability space.

**Examples**

```
method,
bayes_calibration queso
samples = 1000 seed = 348
dram
proposal_covariance
    filename ...
```

```
7.2.68.1.14.9 diagonal
```

- Keywords Area
- method
- bayes_calibration
- queso
- proposal_covariance
- filename
- diagonal

Specifies the diagonal matrix format when importing a user-specified proposal covariance.
Keywords Area

Specification

Alias: none
Argument(s): none

Description

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.

Examples

method,
    bayes_calibration queso
    samples = 1000 seed = 348
    dram
    proposal_covariance
diagonal filename 'dakota_cantilever_queso.diag.dat'

matrix

• Keywords Area
• method
• bayes_calibration
• queso
• proposal_covariance
• filename
• matrix

Specifies the full matrix format when importing a user-specified proposal covariance.

Specification

Alias: none
Argument(s): none

Description

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.

Examples

method,
    bayes_calibration queso
    samples = 1000 seed = 348
    dram
    proposal_covariance
    matrix filename 'dakota_cantilever_queso.matrix.dat'
7.2.68.1.15 options_file

- Keywords Area
- method
- bayes_calibration
- queso
- options_file

File containing advanced QUESO options

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no advanced options file

**Description**

Allow power users to override default QUESO options such as hyper-parameter priors, initial chain position, and proposal covariance (including hyper-parameters), which might not be accessible through Dakota user input.

**Default Behavior** No advanced options used.

**Usage Tips** The options_file offers a final override over other options settings, which are applied with the following precedence:

1. QUESO library defaults
2. Dakota hard-coded defaults
3. Settings from Dakota input file
4. Settings from the advanced options_file

**Examples**

```
# FILE: dakota_queso.opts
# Base QUESO Environment options
env_subDisplayFileName  = QuesoDiagnostics/display
env_subDisplayAllowedSet = 0 1
env_displayVerbosity    = 2
env_seed                 = 2460
```

7.2.68.2 gpmsa

- Keywords Area
- method
- bayes_calibration
- gpmsa

(Experimental) Gaussian Process Models for Simulation Analysis (GPMSA) Bayesian calibration
Topics

This keyword is related to the topics:

- package_queso
- bayesian_calibration

Specification

**Alias:** none  

**Argument(s):** none  

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required</strong></td>
<td></td>
<td>chain_samples</td>
<td>Number of Markov Chain Monte Carlo posterior samples</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>seed</td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>rng</td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td><strong>Required</strong></td>
<td></td>
<td>build_samples</td>
<td>Number of initial model evaluations used in build phase</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>standardized_space</td>
<td>Perform Bayesian inference in standardized probability space</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>logit_transform</td>
<td>Utilize the logit transformation to reduce sample rejection for bounded domains</td>
</tr>
</tbody>
</table>
### Description

GPMSA (Gaussian Process Models for Simulation Analysis) is a surrogate-based Markov Chain Monte Carlo Bayesian calibration method. Dakota’s GPMSA is an experimental capability and not ready for production use at this time.

Central to GPMSA is the construction of a Gaussian Process emulator from simulation runs collected at various settings of input parameters. The emulator is a statistical model of the system response, and it is used to incorporate the observational data to improve system predictions and constrain or calibrate the unknown parameters. The GPMSA code draws heavily on the theory developed in the seminal Bayesian calibration paper by Kennedy and O’Hagan [56]. The particular approach in GPMSA was developed by the Los Alamos National Laboratory statistics group and documented in [49]. Dakota’s GPMSA capability comes from the QUESO package developed at UT Austin.

### Usage Tips:

Configuring GPMSA essentially involves identifying the simulation build data, the experiment data, the calibration and configuration (state) variables, and any necessary algorithm controls. The GP surrogate model is automatically constructed internal to the algorithm and need not be specified through Dakota input.

Dakota’s GPMSA implementation is not intended for production use. There are a number of known limitations, including:

- Only works for scalar and multivariate responses, not field responses. Field responses will be treated as a single multi-variate response set. Consequently, simulation and experiment data must have the same dimensions.
• When build data is not imported a design of experiments will be conducted over all calibration and scenario variables present.

• Experiment data is required (one cannot pose the simulation data as a set of residuals with the assumption of 0-valued experiments).

• Output and diagnostics are limited. Advanced users will need to examine QUESO output files (potentially written in a transformed scaled space) in the QuesoDiagnostic directory.

Examples

The following input file fragment illustrates GPMSA-based Bayesian calibration of $3 \beta$ variables with a uniform prior, with 3 configuration (scenario) variables $x$. A total of 60 simulation build points are provided in `sim_data.dat`, which contains columns for each $\beta$, followed by each $x$, and then the simulation response ‘lin’. Each row of the experiment data file `y_exp_with_var.dat` contains the values of the 3 $x$ variables, followed by the value of ‘lin’ and its observation error (variance).

```
method
  bayes_calibration gpmsa
  chain_samples 1000
  seed 2460
  build_samples 60
  import_build_points_file 'sim_data.dat' freeform
  export_chain_points_file 'posterior.dat'
  burn_in_samples = 100 sub_sampling_period = 2
  posterior_stats kl

variables
  uniform_uncertain 3
    upper_bounds 0.4500 -0.1000 0.4000
    initial_point 0.2750 -0.3000 0.1000
    lower_bounds -0.1000 -0.5000 -0.2000
  descriptors 'beta0' 'beta1' 'beta2'

  continuous_state 3
    upper_bounds 3 * 1.0
    initial_state 3 * 0.5
    lower_bounds 3 * 0.0
  descriptors 'x0' 'x1' 'x2'

responses
  descriptors 'lin'
  calibration_terms 1
  calibration_data_file 'y_exp_with_var.dat'
  freeform
  num_experiments 5
  num_config_variables 3
  experiment_variance_type 'scalar'
  no_gradients
  no_hessians

7.2.68.2.1 chain_samples

  • Keywords Area
  • method
  • bayes_calibration
  • gpmsa
  • chain_samples

Number of Markov Chain Monte Carlo posterior samples
7.2 method

Specification

Alias: samples
Argument(s): INTEGER
Default: method-dependent

Description

The chain_samples keyword indicates the number of draws from the posterior distribution to perform. When an emulator is active, this will be the number of samples on the constructed surrogate model.

Default Behavior

The default number of chain samples is method-dependent. QUESO and MUQ methods use 1000. DREAM uses (number of generations) x (number of chains), resulting in chain_samples close to that specified.

Usage Tips

MCMC methods typically require a large number of chain samples to converge, often thousands to millions.

Examples

```plaintext
method bayes_calibration queso
  chain_samples = 20000
```

7.2.68.2.2 seed

- Keywords Area
- method
- bayes_calibration
- gpmsa
- seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.
Usage Tips

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```plaintext
method
   sampling
      sample_type lhs
      samples = 10
      seed = 15347
```

7.2.68.2.3 `rng`

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmfa**
- **rng**

Selection of a random number generator

Specification

**Alias:** none

**Argument(s):** none

**Default:** Mersenne twister (mt19937)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>RNG Algorithm (Group 1)</td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

**Description**

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.
Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.68.2.3.1 mt19937

- Keywords Area
- method
- bayes_calibration
- gpmsa
- rng
- mt19937

Generates random numbers using the Mersenne twister

Specification

**Alias:** none
**Argument(s):** none

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.68.2.3.2 rnum2

- Keywords Area
- method
- bayes_calibration
- gpmsa
Generates pseudo-random numbers using the Pecos package

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The `rnum2` keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

**Default Behavior**

`rnum2` is not used by default. To change this behavior, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended over `rnum2`.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.68.2.4 build_samples

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gprmsa**
- **build_samples**

Number of initial model evaluations used in build phase

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of `build_samples` is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.
7.2 method

Examples

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```plaintext
method
gpais
    build_samples = 100
    samples_on_emulator = 100
    max_iterations = 5
    response_levels = -1.065
```

7.2.68.2.5 import_build_points_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmsa**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>custom-annotated Tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td></td>
<td></td>
<td>Selects annotated Tabular file format</td>
</tr>
<tr>
<td>freeform</td>
<td></td>
<td></td>
<td>Selects Freeform Tabular file format</td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**
Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method
  polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### 7.2.68.2.5.1 custom_annotated

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmsa**
- **import_build_points_file**
- **custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
**7.2 method**

<table>
<thead>
<tr>
<th>Optional</th>
<th>eval_id</th>
<th>Enable evaluation ID column in custom-annotated tabular file</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
```

7.2.68.2.5.2 header

- Keywords Area
- method
- bayes_calibration
- gpmsa

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• import_build_points_file
• custom_annotated
• header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

7.2.68.2.5.3 eval_id

• Keywords Area
• method
• bayes_calibration
• gpmsa
• import_build_points_file
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.68.2.5.4 interface_id

• Keywords Area
• method
• bayes_calibration
• gpmsa
• import_build_points_file
Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent custom_annotated

Annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

**Usage Tips**
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated` header `eval_id`.

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

**Examples**

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:
```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.2.5.6 `freeform`

- Keywords Area
- method
- bayes_calibration
- gpmsa
- import_build_points_file
- freeform

Selects freeform file format

**Topics**

This keyword is related to the topics:

- `file_formats`

**Specification**

Alias: none

Argument(s): none

Default: annotated format
Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
  freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
```

Perform Bayesian inference in standardized probability space

7.2.68.2.6 standardized_space

• Keywords Area

• method

• bayes_calibration

• gpmsa

• standardized_space

Perform Bayesian inference in standardized probability space

Specification

**Alias**: none

**Argument(s)**: none
Description

This option transforms the inference process (MCMC sampling and any emulator model management) into a standar-
dardized probability space.

The variable transformations performed are as described in askey.

Default Behavior

The default for the Gaussian process and no emulator options is to perform inference in the original probability
space (no transformation). Polynomial chaos and stochastic collocation emulators, on the other hand, are always
formed in standardized probability space, such that the inference process is also performed in this standardized
space.

Expected Output

The user will see the truth model evaluations performed in the original space, whereas any method diagnostics
relating to the MCMC samples (e.g., QUESO data in the outputData directory) will report points and response
data (response gradients and Hessians, if present, will differ but response values will not) that correspond to the
transformed space.

Usage Tips

Selecting standardized_space generally has the effect of scaling the random variables to be of comparable magni-
tude, which can improve the efficiency of the Bayesian inference process.

Examples

method,
  bayes_calibration queso
  samples = 2000 seed = 348
  dram
  standardized_space

7.2.68.2.7 logit_transform

- Keywords Area
- method
- bayes_calibration
- gpmsa
- logit_transform

Utilize the logit transformation to reduce sample rejection for bounded domains

Specification

Alias: none

Argument(s): none

Description

The logit transformation performs an internal variable transformation from bounded domains to unbounded domains
in order to reduce sample rejection due to an out-of-bounds condition.

Default Behavior

This option is experimental at present, and is therefore defaulted off.
Usage Tips
This option can be helpful when regions of high posterior density exist in the corners of a multi-dimensional bounded domain. In these cases, it may be difficult to generate feasible samples from the proposal density, such that transformation to unbounded domains may greatly reduce sample rejection rates.

Examples

```plaintext
method,  
    bayes_calibration queso  
    samples = 2000 seed = 348  
    dram  
    logit_transform
```

7.2.68.2.8 gpmsa_normalize

- Keywords Area
- method
- bayes_calibration
- gpmsa
- gpmsa_normalize

Enable GPMSA-internal normalization

Specification

**Alias:** none
**Argument(s):** none

Description

Normalize variables and data (surrogate model build and experiment) when constructing the underlying Gaussian Process emulator. When specified, this option will normalize calibration parameters and configuration (scenario) variables based on the min/max of the build data. It will normalize simulation and experiment data by the mean and variance of the simulation build data.

**Default Behavior** No GPMSA-specific scaling used.

Examples

```plaintext
method  
    bayes_calibration gpmsa  
    chain_samples 1000  
    seed 2460  
    ...  
    gpmsa_normalize
```

7.2.68.2.9 export_chain_points_file

- Keywords Area
- method
- bayes_calibration
- gpmsa
• **export_chain_points_file**

Export the MCMC chain to the specified filename

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** chain export to default filename

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional (Choose One)</strong></td>
<td><strong>Tabular Format (Group 1)</strong></td>
<td><strong>custom_annotated</strong></td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The filename to which the final MCMC posterior chain will be exported.

**Default Behavior** No export to file.

**Expected Output**

A tabular data file will be produced in the specified format (annotated by default) containing samples from the posterior distribution.

**Usage Tips**

**Additional Discussion**

7.2.68.2.9.1 custom_annotated

• Keywords Area
  • method
  • bayes_calibration
  • gpmsa
  • export_chain_points_file
  • custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

• file_formats
### Specification

**Alias:** none  
**Argument(s):** none

#### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment  
tabular_data  
tabular_data_file = 'dakota_summary.dat'  
custom_annotated header eval_id
```

Resulting tabular file:
\begin{verbatim}
%eval_id   x1     x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9    1.1  0.0002   0.26      0.76
2  0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3  0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...
\end{verbatim}

7.2.68.2.9.2 \texttt{header}

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{bayes\_calibration}
- \texttt{gpmsa}
- \texttt{export\_chain\_points\_file}
- \texttt{custom\_annotated}
- \texttt{header}

Enable header row in custom-annotated tabular file

\textbf{Specification}

\textbf{Alias}: none

\textbf{Argument(s)}: none

\textbf{Description}

See description of parent \texttt{custom\_annotated}

7.2.68.2.9.3 \texttt{eval\_id}

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{bayes\_calibration}
- \texttt{gpmsa}
- \texttt{export\_chain\_points\_file}
- \texttt{custom\_annotated}
- \texttt{eval\_id}

Enable evaluation ID column in custom-annotated tabular file

\textbf{Specification}

\textbf{Alias}: none

\textbf{Argument(s)}: none

\textbf{Description}

See description of parent \texttt{custom\_annotated}
7.2.68.2.9.4  interface_id

• Keywords Area
• method
• bayes_calibration
• gpmsa
• export_chain_points_file
• custom.annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom.annotated

7.2.68.2.9.5  annotated

• Keywords Area
• method
• bayes_calibration
• gpmsa
• export_chain_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.2.9.6 freeform

- Keywords Area
- method
- bayes_calibration
- gpmsa
- export_chain_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
7.2 method

Specification

Alias: none
Argument(s): none
Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
    tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.2.10 dram

- Keywords Area
- method
- bayes_calibration
- gpmsa
- dram

Use the DRAM MCMC algorithm
Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Default: dram

Description

The type of Markov Chain Monte Carlo used. This keyword specifies the use of DRAM, (Delayed Rejection Adaptive Metropolis)[40].

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Usage Tips

If the user knows very little about the proposal covariance, using dram is a recommended strategy. The proposal covariance is adaptively updated, and the delayed rejection may help improve low acceptance rates.

Examples

```plaintext
method,  bayes_calibration queso
dram
samples = 10000 seed = 348
```

7.2.68.2.11  delayed_rejection

- Keywords Area

- method

- bayes_calibration

- gpmsa

- delayed_rejection

Use the Delayed Rejection MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration
7.2 method

Specification

Alias: none
Argument(s): none
Default: dram

Description

This keyword specifies the use of the Delayed Rejection algorithm in which there can be a delay in rejecting samples from the chain. That is, the "DR" part of DRAM is used but the "AM" part is not, rather a regular Metropolis-Hastings algorithm is used.

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Usage Tips

If the user knows something about the proposal covariance or the proposal covariance is informed through derivative information, using delayed_rejection is preferred over dram: the proposal covariance is already being informed by derivative information and the adaptive metropolis is not necessary.

Examples

```
method,
    bayes_calibration queso
delayed_rejection
    samples = 10000 seed = 348
```

See Also

These keywords may also be of interest:

- proposal_covariance

7.2.68.2.12 adaptive_metropolis

- Keywords Area
- method
- bayes_calibration
- gpmsa
- adaptive_metropolis

Use the Adaptive Metropolis MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration
Specification

**Alias:** none

**Argument(s):** none

**Default:** dram

**Description**

This keyword specifies the use of the Adaptive Metropolis algorithm. That is, the "AM" part of DRAM is used but the "DR" part is not: specifying this keyword activates only the Adaptive Metropolis part of the MCMC algorithm, in which the covariance of the proposal density is updated adaptively.

**Default Behavior**

Five MCMC algorithm variants are supported in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Four MCMC variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings and adaptive_metropolis.

**Usage Tips**

If the user knows very little about the proposal covariance, but doesn’t want to incur the cost of using full dram with both delayed rejection and adaptive metropolis, specifying only adaptive_metropolis offers a good strategy.

**Examples**

```
method,
    bayes_calibration queso
    adaptive_metropolis
    samples = 10000 seed = 348
```

**7.2.68.2.13 metropolis_hastings**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmsa**
- **metropolis_hastings**

Use the Metropolis-Hastings MCMC algorithm

**Topics**

This keyword is related to the topics:

- **bayesian_calibration**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** dram
Description

This keyword specifies the use of a Metropolis-Hastings algorithm for the MCMC chain generation. This means there is no delayed rejection and no adaptive proposal covariance updating as in DRAM.

Default Behavior

Five MCMC algorithm variants are supported currently in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Four MCMC algorithm variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings and adaptive_metropolis.

Usage Tips

If the user wants to use Metropolis-Hastings, possibly as a comparison to the other methods which involve more chain adaptation, this is the MCMC type to use.

Examples

```plaintext
method,
  bayes_calibration queso
  metropolis_hastings
  samples = 10000 seed = 348
```

7.2.68.2.14 proposal_covariance

- Keywords Area
- method
- bayes_calibration
- gpmsa
- proposal_covariance

Defines the technique used to generate the MCMC proposal covariance.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

- Alias: none
- Argument(s): none
- Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Proposal Covariance Source (Group 1)</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>prior</td>
<td>Uses the covariance of the prior distributions to define the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>derivatives</td>
<td>Use derivatives to inform the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>values</td>
<td>Specifies matrix values to use as the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>filename</td>
<td>Uses a file to import a user-specified MCMC proposal covariance.</td>
</tr>
</tbody>
</table>

**Description**

The proposal covariance is used to define a multivariate normal (MVN) jumping distribution used to create new points within a Markov chain. That is, a new point in the chain is determined by sampling within a MVN probability density with prescribed covariance that is centered at the current chain point. The accuracy of the proposal covariance has a significant effect on rejection rates and the efficiency of chain mixing.

**Default Behavior**

The default proposal covariance is prior when no emulator is present; derivatives when an emulator is present.

**Expected Output**

The effect of the proposal covariance is reflected in the MCMC chain values and the rejection rates, which can be seen in the diagnostic outputs from the QUESO solver within the QuesoDiagnostics directory.

**Usage Tips**

When derivative information is available inexpensively (e.g., from an emulator model), the derived-based proposal covariance forms a more accurate proposal distribution, resulting in lower rejection rates and faster chain mixing.

**7.2.68.2.14.1 prior**

- Keywords Area
- method
- bayes_calibration
- gpmsa
- proposal_covariance
- prior

Uses the covariance of the prior distributions to define the MCMC proposal covariance.
Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>multiplier</td>
<td>Multiplier to scale prior variance</td>
</tr>
</tbody>
</table>

Description

This keyword selection results in definition of the MCMC proposal covariance from the covariance of the prior distributions. This covariance is currently assumed to be diagonal without correlation.

Default Behavior

This is the default proposal_covariance option.

Usage Tips

Since this proposal covariance is defined globally, the chain does not need to be periodically restarted using local updates to this proposal. However, it is usually effective to adapt the proposal using one of the adaptive metropolis MCMC options.

Examples

```plaintext
method, bayes_calibration queso
   samples = 2000 seed = 348
   dram
   proposal_covariance prior
```

7.2.68.2.14.2  **multiplier**

- Keywords Area
- method
- bayes_calibration
- gpmsa
- proposal_covariance
- prior
- multiplier

Multiplier to scale prior variance
Specification

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0

**Description**

The initial proposal covariance will be given by the prior variance times the multiplier.

**Default Behavior** When using prior-based proposal covariance, the default is to use the prior variance for the proposal (multiplier = 1.0)

**Usage Tips**

The prior variance may result in too tight or narrow a proposal covariance. The multiplier can be used to scale all entries of the prior variance in determining the initial proposal covariance.

**Examples**

```plaintext
method bayes_calibration queso
   samples = 2000 seed = 348
dram
   proposal_covariance prior
   multiplier 0.1
```

**7.2.68.2.14.3 derivatives**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmsa**
- **proposal_covariance**
- **derivatives**

Use derivatives to inform the MCMC proposal covariance.

**Topics**

This keyword is related to the topics:

- **bayesian_calibration**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**
### Description

This keyword selection results in definition of the MCMC proposal covariance from the Hessian of the misfit function (negative log likelihood), where this Hessian is defined from either a Gauss-Newton approximation (using only first derivatives of the calibration terms) or a full Hessian (using values, first derivatives, and second derivatives of the calibration terms). If this Hessian is indeterminate, it will be corrected as described in [16].

#### Default Behavior

The default is prior based proposal covariance. This is a more advanced option that exploits structure in the form of the likelihood.

#### Expected Output

When derivatives are specified for defining the proposal covariance, the misfit Hessian and its inverse (the MVN proposal covariance) will be output to the standard output stream.

#### Usage Tips

The full Hessian of the misfit is used when either supported by the emulator in use (for PCE and surfpack GP, but not SC or dakota GP) or by the user's response specification (Hessian type is not "no_hessians"), in the case of no emulator. When this full Hessian is indefinite and cannot be inverted to form the proposal covariance, fallback to the positive semi-definite Gauss-Newton Hessian is employed.

Since this proposal covariance is locally accurate, it should be updated periodically using the `update_period` option. While the adaptive metropolis option can be used in combination with derivative-based preconditioning, it is generally preferable to instead decrease the proposal update period due to the improved local accuracy of this approach.

#### Examples

Generate a 2000 sample posterior chain, using derivatives to initialize the proposal covariance at the start of the chain.

```plaintext
method,
bayes_calibration queso
  samples = 2000 seed = 348
delayed_rejection
  emulator_pce sparse_grid_level = 2
  proposal_covariance derivatives

7.2.68.2.14.4 update_period
  • Keywords Area
  • method
  • bayes_calibration
  • gpmsa
  • proposal_covariance
  • derivatives
  • update_period
```

Period at which to update derivative-based proposal covariance
Specification

Alias: none
Argument(s): INTEGER

Description

For derivative-based proposal covariance, this specifies the period (number of accepted MCMC samples) after which the proposal covariance is updated using derivative values at the current chain point.

Default Behavior

When update_period is not specified, derivatives will inform the proposal covariance at the start of the chain, but not updated further.

Usage Tips

The update_period should be tailored to the size of the total chain, accounting for the relative expense of derivative-based proposal updates.

Examples

method,
   bayes_calibration queso
   samples = 2000 seed = 348
   delayed_rejection
   emulator pce sparse_grid_level = 2
   proposal_covariance derivatives
      update_period = 40  # update proposal covariance every 40 points

7.2.68.2.14.5 values

  • Keywords Area
  • method
  • bayes_calibration
  • gpmsa
  • proposal_covariance
  • values

Specifies matrix values to use as the MCMC proposal covariance.

Specification

Alias: none
Argument(s): REALLIST
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Values For (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required</strong> (Choose One)</td>
<td></td>
<td>diagonal</td>
<td>Specifies the diagonal matrix format when specifying a user-specified proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>matrix</td>
<td>Specifies the full matrix format when specifying a user-specified proposal covariance.</td>
</tr>
</tbody>
</table>

**Description**

This keyword selection results in definition of the MCMC proposal covariance from user-specified matrix values. The matrix input format must be declared as either a full matrix or a matrix diagonal.

**Default Behavior**

This option is not the default, and generally implies special a priori knowledge from the user.

**Usage Tips**

This option is not supported for the case of transformations to standardized probability space.

**Examples**

```plaintext
method,
    bayes_calibration queso
    samples = 1000 seed = 348
dram
    proposal_covariance
        values ... # See leaf nodes for required format option
```

**7.2.68.2.14.6 diagonal**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **gpmsa**
- **proposal_covariance**
- **values**
- **diagonal**

Specifies the diagonal matrix format when specifying a user-specified proposal covariance.

**Specification**

**Alias:** none

**Argument(s):** none
Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.

Examples

```
method, bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
diagonal values 1.0e6 1.0e-1
```

7.2.68.2.14.7 matrix

- Keywords Area
- method
- bayes_calibration
- gpmsa
- proposal_covariance
- values
- matrix

Specifies the full matrix format when specifying a user-specified proposal covariance.

Specification

Alias: none

Argument(s): none

Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.

Examples

```
method, bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
matrix values 1.0 0.1
  0.1 2.0
```

7.2.68.2.14.8 filename

- Keywords Area
- method
- bayes_calibration
Uses a file to import a user-specified MCMC proposal covariance.

**Specification**

**Alias:** none  
**Argument(s):** STRING

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Values For (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>diagonal</td>
<td>matrix</td>
<td>Specifies the full matrix format when importing a user-specified proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Specifies the diagonal matrix format when importing a user-specified proposal covariance.</td>
</tr>
</tbody>
</table>

**Description**

This keyword selection results in definition of the MCMC proposal covariance from importing data a user-specified filename. This import must be declared as either a full matrix or a matrix diagonal.

**Default Behavior**

This option is not the default, and generally implies special a priori knowledge from the user.

**Usage Tips**

This option is not supported for the case of transformations to standardized probability space.

**Examples**

```bash
method, bayes_calibration queso
samples = 1000 seed = 348
dram
proposal_covariance
filename ... # See leaf nodes for required format option
```

7.2.68.2.14.9 diagonal

- Keywords Area
- method
Specifies the diagonal matrix format when importing a user-specified proposal covariance.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.

**Examples**

```plaintext
method,
  bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
    diagonal filename 'dakota_cantilever_queso.diag.dat'
```

Specifications the full matrix format when importing a user-specified proposal covariance.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.
Examples

```method,
bayes_calibration queso
  samples = 1000 seed = 348
dram
  proposal_covariance
    matrix filename 'dakota_cantilever_queso.matrix.dat'
```

### 7.2.68.2.15 options_file

- **Keywords**
  - Area
  - method
  - bayes_calibration
  - gpmsa
  - options_file

File containing advanced QUESO/GPMSA options

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no advanced options file

**Description**

Allow power users to override default QUESO/GPMSA options such as hyper-parameter priors, initial chain position, and proposal covariance (including hyper-parameters), which might not be accessible through Dakota user input.

**Default Behavior** No advanced options used.

**Usage Tips** The `options_file` offers a final override over other options settings, which are applied with the following precedence:

1. QUESO library defaults
2. Dakota hard-coded defaults
3. Settings from Dakota input file
4. Settings from the advanced `options_file`

**Examples**

```bash
# FILE: dakota_gpmsa.opts

# Base QUESO Environment options
env_subDisplayFileName = QuesoDiagnostics/display
env_subDisplayAllowedSet = 0 1
env_displayVerbosity = 2
env_seed = 2460

# GPMSA Options
gpmsa_emulator_precision_shape = 2.0
gpmsa_emulator_precision_scale = 0.2
gpmsa_emulator_correlation_strength_alpha = 1.0
```
gpmsa_emulator_correlation_strength_beta = 0.1
gpmsa_discrepancy_precision_shape = 1.0
gpmsa_discrepancy_precision_scale = 1e4
gpmsa_discrepancy_correlation_strength_alpha = 1.0
gpmsa_discrepancy_correlation_strength_beta = 0.1
gpmsa_emulator_data_precision_shape = 3.0
gpmsa_emulator_data_precision_scale = 333.333

7.2.68.3 wasabi

  • Keywords Area

  • method

  • bayes_calibration

  • wasabi

(Experimental Method) Non-MCMC Bayesian inference using interval analysis

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>pushforward_-samples</td>
<td></td>
<td>(Experimental Capability) Number of samples of the prior to push forward through the model. Description: WASABI requires a forward UQ that maps samples from the prior parameter distribution through the model. The corresponding responses then are used to see their relative likelihood according to the density of the observational data. The <code>pushforward_-samples</code> refers to the number of samples taken from the prior and used in the forward UQ. It typically should be high, e.g., 10000, of which only a small fraction may be in a high density region of the posterior.</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>emulator</td>
<td></td>
<td>Use an emulator or surrogate model to evaluate the likelihood function</td>
</tr>
<tr>
<td>Optional</td>
<td>standardized_-space</td>
<td></td>
<td>Perform Bayesian inference in standardized probability space</td>
</tr>
<tr>
<td>Type</td>
<td>Keyword</td>
<td>Description</td>
<td></td>
</tr>
<tr>
<td>-----------</td>
<td>--------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Required</td>
<td>data_distribution</td>
<td>(Experimental Capability) Specify the distribution of the experimental data</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>posterior_samples-import_filename</td>
<td>Filename for samples at which the user would like the posterior density calculated</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>generate-posterior_samples</td>
<td>(Experimental Capability) Generate random samples from the posterior density</td>
<td></td>
</tr>
</tbody>
</table>

Description: This keyword will result in samples from the prior that have a high posterior density being printed to a file called 'psamples.txt' as a default, or to a file specified by the posterior-samples-export-filename.
Description

Offers an alternative to Markov Chain Monte Carlo-based Bayesian inference. This is a nascent capability, not yet ready for production use.

Usage Guidelines: The WASABI method requires an emulator model.

Attention: While the emulator specification for WASABI includes the keyword posterior_adaptive, it is not yet operational.

Examples

```
method
   bayes_calibration
      wasabi
```

7.2.68.3.1 pushforward_samples

- Keywords Area
- method
- bayes_calibration
- wasabi
- pushforward_samples

(Experimental Capability) Number of samples of the prior to push forward through the model

Description: WASABI requires a forward UQ that maps samples from the prior parameter distribution through the model. The corresponding responses then are used to see their relative likelihood according to the density of the
observational data. The \texttt{pushforward_samples} refers to the number of samples taken from the prior and used in the forward UQ. It typically should be high, e.g. 10000, of which only a small fraction may be in a high density region of the posterior.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** method-dependent

7.2.68.3.2 seed

- Keywords Area
- method
- bayes_calibration
- wasabi
- seed

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If \texttt{seed} is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without \texttt{seed} specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```
Use an emulator or surrogate model to evaluate the likelihood function

**Specification**

*Alias*: none  
*Argument(s)*: none  
*Child Keywords*:

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emulator Type (Group 1)</td>
<td></td>
<td>gaussian_process</td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td>pce</td>
<td></td>
<td></td>
<td>Polynomial Chaos Expansion surrogate model</td>
</tr>
<tr>
<td>ml_pce</td>
<td></td>
<td></td>
<td>Multilevel Polynomial Chaos Expansion as an emulator model.</td>
</tr>
<tr>
<td>mf_pce</td>
<td></td>
<td></td>
<td>Multifidelity Polynomial Chaos Expansion as an emulator model.</td>
</tr>
<tr>
<td>sc</td>
<td></td>
<td></td>
<td>Stochastic Collocation as an emulator model.</td>
</tr>
<tr>
<td>mf_sc</td>
<td></td>
<td></td>
<td>Multifidelity Stochastic Collocation as an emulator model.</td>
</tr>
</tbody>
</table>

**Description**

This keyword describes the type of emulator used when calculating the likelihood function for the Bayesian calibration. The emulator can be a Gaussian process, polynomial chaos expansion, or stochastic collocation.

7.2.68.3.1  gaussian_process

*Keywords Area*
- method
- bayes_calibration
- wasabi
- emulator
- gaussian_process

Gaussian Process surrogate model

Specification

**Alias:** krige

**Argument(s):** none

**Default:** Surfpack Gaussian process

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group GP Implementation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>dakota</td>
<td>Select the built-in Gaussian Process surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>build_samples</td>
<td></td>
<td>Number of initial model evaluations used in build phase</td>
</tr>
<tr>
<td>Optional</td>
<td>posterior_adaptive</td>
<td></td>
<td>Adapt emulator model to increase accuracy in high posterior probability regions</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build_points_file</td>
<td></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

**Description**

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.
Use the Surfpack version of Gaussian Process surrogates

**Specification**

**Alias:** none
**Argument(s):** none

**Description**

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. **Optimization methods:**
   
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See `optimization_method`.
   
   The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. **Trend Function:**
   
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See `trend`.

3. **Correlation Lengths:**
   
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See `correlation_lengths`.

4. **Ill-conditioning**
   
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   
   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolation GP model would become inept at extrapolation and interpolation.
   
   The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using `surfpack`.

5. **Gradient Enhanced Kriging (GEK).**
   
   The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   
   See notes in the Theory section.
Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK’s correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging’s correlation matrix (which is a small sub-matrix within GEK’s correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK’s correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging’s correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK’s correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK’s correlation matrix.

7.2.68.3.3.3 dakota

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- gaussian_process
- dakota

Select the built in Gaussian Process surrogate

Specification

Alias: none
Argument(s): none

Description

A second version of GP surrogates was available in prior versions of Dakota. For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in[60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to
select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

7.2.68.3.3.4 build_samples

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- gaussian_process
- build_samples

Number of initial model evaluations used in build phase

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of build_samples is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

**Examples**

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```plaintext
method
gpais
    build_samples = 100
    samples_on_emulator = 100
    max_iterations = 5
    response_levels = -1.065
```

7.2.68.3.3.5 posterior_adaptive

- Keywords Area
- method
- bayes_calibration
- wasabi
Adapt emulator model to increase accuracy in high posterior probability regions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

**Examples**

```plaintext
bayes_calibration queso
    chain_samples = 2000 seed = 348
delayed_rejection
emulator
gaussian_process surfpack build_samples = 30
posterior_adaptive max_iterations = 10
proposal_covariance derivatives
```

**7.2.68.3.3.6 import_build_points_file**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **gaussian_process**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

#### Default Behavior

By default, methods do not import points from a file.

#### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- `annotated` (default)
- `custom_annotated`
- `freeform`

#### Examples

```
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

```
7.2.68.3.3.7 custom_annotated
```

- Keywords Area
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - gaussian_process
  - import_build_points_file

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
**custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.3.3.8 header

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **gaussian_process**
- **import_build_points_file**
- **custom_annotated**
- **header**

Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no header

**Description**

See description of parent **custom_annotated**

7.2.68.3.3.9 eval_id

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom.annotated

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom.annotated
7.2.6.3.3.11 annotated

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- gaussian_process
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.3.3.12 freeform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- gaussian_process
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips
Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

**Resulting tabular file:**

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1 0.0001996404857  0.2601620081  0.759955
0.89991 1.1 0.0002003604863  0.2598380081  0.760045
...```

**7.2.68.3.3.13 active_only**

- **Keywords Area**
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - gaussian_process
  - import_build_points_file
  - active_only

Import only active variables from tabular data file

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none
Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.3.14 pce

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce

Polynomial Chaos Expansion surrogate model

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_-iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
</tr>
<tr>
<td><strong>Optional (Choose One)</strong></td>
<td><strong>Basis Polynomial Family (Group 2)</strong></td>
<td><strong>askey</strong></td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td>--------------------------</td>
<td>---------------------------------------</td>
<td>-----------</td>
<td>---------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>wiener</strong></td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td><strong>normalized</strong></td>
<td>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials. Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>export_expansion_file</strong></td>
<td></td>
</tr>
</tbody>
</table>
### Description

Selects a polynomial chaos expansion (PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using PCE as a surrogate within the Bayesian framework.

### See Also

These keywords may also be of interest:

- `polynomial_chaos`

#### 7.2.68.3.3.15  p_refinement

- `Keywords Area`
- `method`
- `bayes_calibration`
- `wasabi`
- `emulator`
- `pce`
- `p_refinement`

Automatic polynomial order refinement

### Specification

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group p-refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>diagonal covariance</td>
<td>full_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td></td>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

**Required (Choose One):**

- `diagonal covariance`
- `full covariance`

**Display only the diagonal terms of the covariance matrix**

<table>
<thead>
<tr>
<th>Display only the diagonal terms of the covariance matrix</th>
<th>Display the full covariance matrix</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.3.3.16 uniform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- `p_refinement`
- uniform

Refine an expansion uniformly in all dimensions.

Specification

**Alias**: none
**Argument(s)**: none
The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.3.3.17 dimension_adaptive

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.</td>
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<tr>
<td></td>
<td>decay</td>
<td>Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.</td>
<td></td>
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<tr>
<td>method</td>
<td>description</td>
<td>7.2 method</td>
<td>generalized</td>
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</tr>
<tr>
<td></td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.3.3.18 sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized

### Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.

**Examples**

```
method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```dakota
method,  
  polynomial_chaos  
  sparse_grid_level = 3  
  dimension_adaptive p_refinement generalized  
  max_iterations = 20  
  convergence_tol = 1.e-4
```

**7.2.68.3.3.21 max_refinement_iterations**

- **Keywords** Area  
- **method**  
- **bayes_calibration**  
- **wasabi**  
- **emulator**  
- **pce**  
- **max_refinement_iterations**

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100
Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

method,
    polynomial_chaos
dimension_adaptive generalized
    p_refinement
    max_refinement_iterations = 20
    convergence_tol = 1.e-4
    sparse_grid_level = 1

7.2.68.3.3.22 quadrature_order

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• pce
• quadrature_order

Order for tensor-products of Gaussian quadrature rules

Specification

Alias: none
Argument(s): INTEGER
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional/Choose One</td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order, and, optionally, dimension_preference). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for...
gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of \textit{p\_refinement} or the case of an explicit \textit{nested} override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a \textit{dimension\_preference}, where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified \textit{quadrature\_order} and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order \( p_i \) of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision \( 2m_i - 1 \), \( p_i \) is one less than the quadrature order \( m_i \) in each dimension (a one-dimensional expansion contains the same number of terms, \( p + 1 \), as the number of Gauss points). The total number of terms, \( N \), in a tensor-product expansion involving \( n \) uncertain input variables is

\[
N = 1 + P = \prod_{i=1}^{n} (p_i + 1)
\]

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the \textit{quadrature\_order} specification supports an array input.

A corresponding sequence specification is documented at, e.g., \textit{quadrature\_order\_sequence} and \textit{quadrature\_order\_sequence}

See Also

These keywords may also be of interest:

- \textit{quadrature\_order\_sequence} and \textit{method-multifidelity\_stoch\_collocation-quadrature\_order\_sequence}

7.2.68.3.3.23 \hspace{1em} \textit{dimension\_preference}

- Keywords Area
- method
- bayes\_calibration
- wasabi
- emulator
- pce
- \textit{quadrature\_order}
- \textit{dimension\_preference}

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: REALLIST

\textbf{Default}: isotropic grids

\textbf{Description}

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.3.24 nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- quadrature_order
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.3.25 non_nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules
7.2 method

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.26 sparse_grid_level

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>dimension_—preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 1)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
</tbody>
</table>
Overview

The Smolyak sparse grid method, specified with `sparse_grid_level` and, optionally, `dimension_preference`, allows for multi-dimensional integration. The underlying one-dimensional integration rules are the same as for tensor-product quadrature, but the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). For tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User's Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User's Manual for additional details). As for `quadrature_order`, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`.

See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence

7.2.68.3.3.27 `dimension_preference`

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.3.28 restricted

**Keywords Area**

- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

Specification

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)

**Description**

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid ( \( i = 4l + 1 \) ) or an order \( m \) for a tensor grid ( \( i = 2m - 1 \) ). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it
cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.29 unrestricted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none
Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid ( \( i = 4l + 1 \) ) or an order \( m \) for a tensor grid ( \( i = 2m - 1 \) ). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.30 nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level
7.2 method

- nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.3.31 non_nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- sparse_grid_level
- non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.32 cubature_integrand

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
Cubature using Stroud rules and their extensions

Specification

Alias: none
Argument(s): INTEGER

Description

Multi-dimensional integration by Stroud cubature rules [80] and extensions [94], as specified with cubature_integrand. A total-order expansion is used, where the isotropic order $p$ of the expansion is half of the integrand order, rounded down. The total number of terms $N$ for an isotropic total-order expansion of order $p$ over $n$ variables is given by

$$N = 1 + P = 1 + \sum_{s=1}^{p} \frac{1}{s!} \prod_{r=0}^{s-1} (n + r) = \frac{(n + p)!}{n! p!}$$

Since the maximum integrand order is currently five for normal and uniform and two for all other types, at most second- and first-order expansions, respectively, will be used. As a result, cubature is primarily useful for global sensitivity analysis, where the Sobol' indices will provide main effects and, at most, two-way interactions. In addition, the random variable set must be independent and identically distributed (iid), so the use of askey or wiener transformations may be required to create iid variable sets in the transformed space (as well as to allow usage of the higher order cubature rules for normal and uniform). Note that global sensitivity analysis often assumes uniform bounded regions, rather than precise probability distributions, so the iid restriction would not be problematic in that case.

7.2.68.3.3.33 expansion_order

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order

The (initial) order of a polynomial expansion

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

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<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
### Description

When the `expansion_order` for a polynomial chaos expansion is specified, the coefficients may be computed by integration based on random samples or by regression using either random or sub-sampled tensor product quadrature points.

Multidimensional integration by Latin hypercube sampling (specified with `expansion_samples`). In this case, the expansion order \( p \) cannot be inferred from the numerical integration specification and it is necessary to provide an `expansion_order` to specify \( p \) for a total-order expansion.

Linear regression (specified with either `collocation_points` or `collocation_ratio`). A total-order expansion is used and must be specified using `expansion_order` as described in the previous option. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the `collocation_ratio` allows for specification of a constant factor applied to \( N \) (e.g., `collocation_ratio = 2` produces samples = \( 2N \)). In addition, the default linear

<table>
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<th><code>dimension_preference</code></th>
<th>A set of weights specifying the relative importance of each uncertain variable (dimension)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td><code>basis_type</code></td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td><code>collocation_points</code></td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td></td>
<td><code>collocation_ratio</code></td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td></td>
<td><code>expansion_samples</code></td>
<td>Number of simulation samples used to estimate the expected value of a set of PCE coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td><code>import_build_points_file</code></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td><code>posterior_adaptive</code></td>
<td>Adapt emulator model to increase accuracy in high posterior probability regions</td>
</tr>
</tbody>
</table>
relationship with $N$ can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $\frac{cN^o}{r+1}$). When admissible, a constrained least squares approach is employed in which response values are first reproduced exactly and error in reproducing response derivatives is minimized. Two collocation grid options are supported: the default is Latin hypercube sampling ("point collocation"), and an alternate approach of "probabilistic collocation" is also available through inclusion of the tensor_grid keyword. In this alternate case, the collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

If collocation_points or collocation_ratio is specified, the PCE coefficients will be determined by regression. If no regression specification is provided, appropriate defaults are defined. Specifically SVD-based least-squares will be used for solving over-determined systems and under-determined systems will be solved using LASSO. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares. Technical information on the various methods listed below can be found in the Linear regression section of the Theory Manual. Some of the regression methods (O-MP, LASSO, and LARS) are able to produce a set of possible PCE coefficient vectors (see the Linear regression section in the Theory Manual). If cross validation is inactive, then only one solution, consistent with the noise_tolerance, will be returned. If cross validation is active, Dakota will choose between possible coefficient vectors found internally by the regression method across the set of expansion orders (1,....,expansion_order) and the set of specified noise tolerances and return the one with the lowest cross validation error indicator.

7.2.68.3.34 dimension_preference

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none

Argument(s): REALLIST

Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional Description of Group</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong> tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td><strong>PCE Basis Type (Group 1)</strong> total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td>adapted</td>
</tr>
</tbody>
</table>

**Description**

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order \( p \) keep all terms with \( d \)-dimensional multi index \( i = (i_1, \ldots, i_d) \) that satisfies

\[
\max(i_1, \ldots, i_d) \leq p
\]

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none
Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.68.3.3.38 adapted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>advancements</td>
<td>The maximum number of steps used to expand a basis step. The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>soft_convergence_limit</td>
<td></td>
</tr>
</tbody>
</table>

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.3.3.39  advancements

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

Specification

**Alias:** none

**Argument(s):** INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.40  soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.
7.2 method

Specification

Alias: none
Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.41 collocation_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points

Number of collocation points used to estimate expansion coefficients

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td><strong>Optional</strong> (Choose One)</td>
<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
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<td>Keywords Area</td>
<td>Description</td>
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<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
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<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
<td></td>
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<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Optional | **cross_validation** | Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion. |
| Optional | **ratio_order** | Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients. |
| Optional | **response_scaling** | Perform bounds-scaling on response values prior to surrogate emulation. |
| Optional | **use_derivatives** | Use derivative data to construct surrogate models. |
| Optional | **tensor_grid** | Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion. |
| Optional | **reuse_points** | This describes the behavior of reuse of points in constructing polynomial chaos expansion models. |
Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches. A corresponding sequence specification is documented at, e.g., collocation_points_sequence and collocation_points_sequence.

See Also

These keywords may also be of interest:

- collocation_points_sequence method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence

7.2.68.3.3.42 least_squares

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

Alias: none

Argument(s): none

Default: svd

Child Keywords:

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<tr>
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<td>LSQ Regression Approach (Group 1)</td>
<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
</tr>
<tr>
<td>equality_constrained</td>
<td></td>
<td></td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.68.3.3.43 svd

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_squares
- svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Specification

**Alias**: none

**Argument(s)**: none

Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

7.2.68.3.3.44 equality_constrained

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_squares
- equality_constrained

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.
7.2 method

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.3.45  orthogonal_matching_pursuit

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• pce
• expansion_order
• collocation_points
• orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.3.46  noise_tolerance

• Keywords Area
• method
• bayes_calibration
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.47  basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp  
**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
7.2.68.3.3.48  basis_pursuit_denoising

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

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<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.3.3.49  noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- basis_pursuit_denoising
• noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.50 least_angle_regression

• Keywords Area

• method

• bayes_calibration

• wasabi

• emulator

• pce

• expansion_order

• collocation_points

• least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:

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<tr>
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<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
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</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
7.2 method

7.2.68.3.3.51 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.52 least_absolute_shrinkage

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso

Argument(s): none

Child Keywords:
## 7.2.68.3.53 noise_tolerance

Compute the coefficients of a polynomial expansion by using the LASSO problem.

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### Specification

**Alias:** none  
**Argument(s):** REALIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
7.2.6.3.3.54 l2_penalty

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

Description

The $l_2$ penalty used when performing compressed sensing with elastic net.

7.2.6.3.3.55 cross_validation

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.3.3.56 noise_only

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** false

### Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.
7.2.68.3.3.57 max_cv_order_candidates

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

**Description**

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

**See Also**

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train
Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

**Description**

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \(cN^{o}\) where \(c\) is the collocation_ratio and \(o\) is the ratio_order.

**Perform bounds-scaling on response values prior to surrogate emulation**
7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Default**: use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.68.3.60 use_derivatives

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_points**
- **use_derivatives**

Use derivative data to construct surrogate models

Specification

**Alias**: none

**Argument(s)**: none

**Default**: use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it's use with Surfpack Gaussian process is not recommended.
Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
7.2 method

Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The \textit{reuse_points} option controls the reuse behavior of points for various types of polynomial chaos expansions, including: \textit{collocation_points}, \textit{collocation_ratio}, \textit{expansion_samples}, or \textit{orthogonal_least_interpolation}. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify \textit{reuse_points} so that any points that have been previously generated (for example, from the \textit{import_points} file) can be reused.

7.2.68.3.3.63 max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_points
- max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.3.3.64 collocation_ratio

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Child Keywords:**

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<tr>
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<th>Regression Algorithm (Group 1)</th>
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<tr>
<td>Optional</td>
<td></td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td>Method</td>
<td>Description</td>
<td></td>
<td></td>
</tr>
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<td>-----------------------------</td>
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<tr>
<td><code>least_angle_regression</code></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>least_absolute_shrinkage</code></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional cross_validation</td>
<td>Use cross validation to choose the ‘best’ polynomial order of a polynomial chaos expansion.</td>
<td></td>
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<tr>
<td>Optional ratio_order</td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients. Perform bounds-scaling on response values prior to surrogate emulation.</td>
<td></td>
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</tr>
<tr>
<td>Optional response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation.</td>
<td></td>
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</tr>
<tr>
<td>Optional use_derivatives</td>
<td>Use derivative data to construct surrogate models.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o + 1 \)).

7.2.68.3.3.65 least_squares

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

**Alias**: none

**Argument(s)**: none

**Default**: svd

**Child Keywords**:

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<tr>
<td>Optional/Choose One</td>
<td></td>
<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.6.8.3.3.66 svd

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- least_squares
- svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Specification

Alias: none

Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

7.2.6.8.3.3.67 equality_constrained

- Keywords Area
- method
- bayes_calibration
- wasabi
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.3.3.68    orthogonal_matching_pursuit

- **Keywords Area**
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.3.3.69 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.70 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
Specification

**Alias:** bp

**Argument(s):** none

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.3.3.71 basis_pursuit_denoising

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

Specification

**Alias:** bpdn

**Argument(s):** none

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.
7.2.68.3.3.72 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.73 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:
**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.3.3.74  noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

*Alias*: none  
*Argument(s)*: REALLIST  
*Default*: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.75  least_absolute_shrinkage

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
7.2 method

- expansion_order
- collocation_ratio
- least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>l2_penalty</td>
<td>l2_penalty</td>
<td></td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.3.3.76 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.77  l2_penalty

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_ratio**
- **least_absolute_shrinkage**
- **l2_penalty**

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.

7.2.68.3.78  cross_validation

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **expansion_order**
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

**7.2.68.3.3.79 noise_only**

- **Keywords Area**
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - pce
  - expansion_order
  - collocation_ratio
  - cross_validation
  - noise_only

Restrict the cross validation process to estimating only the best noise tolerance.
Specification

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.3.80 max_cv_order_candidates

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_ratio**
- **cross_validation**
- **max_cv_order_candidates**

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

**Description**

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.
See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.3.3.81 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none
Argument(s): REAL
Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to \([0,1]\) based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- multifidelity_polynomial_chaos
- multifidelity_function_train

---

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Use derivative data to construct surrogate models

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it's use with Surfpack Gaussian process is not recommended.

7.2.68.3.84 tensor_grid

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
7.2.68.3.3.85  reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

### Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

### Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

7.2.68.3.3.86  max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients
Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.3.3.87 expansion_samples

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- expansion_samples

Number of simulation samples used to estimate the expected value of a set of PCE coefficients

Specification

Alias: none
Argument(s): INTEGER
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

Description

In this case, the expansion order $p$ cannot be inferred from the numerical integration specification and it is necessary to provide an expansion_order to specify $p$ for a total-order expansion.

A corresponding sequence specification is documented at, e.g., expansion_samples_sequence and expansion_samples_sequence
See Also

These keywords may also be of interest:

- `expansion_samples_sequence`
- `method-multilevel_polynomial_chaos-expansion_order_sequence-expansion_samples_sequence`  

7.2.68.3.3.88 reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- expansion_samples
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.68.3.3.89 import_build_points_file

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- expansion_order
- import_build_points_file

File containing points you wish to use to build a surrogate
Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated *(default)*
- custom_annotated
- freeform

**Examples**

```plaintext
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.3.90 custom_annotated

- Keywords Area
- method
Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.
### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1    x2    obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9    1.1    0.0002  0.26    0.76
2  0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3  0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...  
```

### 7.2.68.3.3.91 header

- **Keywords Area**
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - pce
  - expansion_order
  - import_build_points_file
  - custom_annotated
  - header

Enable header row in custom-annotated tabular file

### Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

### Description

See description of parent custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

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Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.3.3.94 annotated

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• pce
• expansion_order
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Selects freeform file format

Comparison

Selects freeform file format

Topics

This keyword is related to the topics:

- `file_formats`
7.2 method

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1  0.0001996404857  0.2601620081  0.759955
0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.3.3.96 active_only

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
Import only active variables from tabular data file

Topics
This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.3.97 posterior_adaptive

Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none
Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

Examples

bayes_calibration queso
  chain_samples = 2000 seed = 348
delayed_rejection
evaluator
gaussian_process surfpack build_samples = 30
posterior_adaptive max_iterations = 10
proposal_covariance derivatives

7.2.68.3.3.98 orthogonal_least_interpolation

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>collocation_points</td>
<td></td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>
### Optional

<table>
<thead>
<tr>
<th>Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>reuse_points</code></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models. File containing points you wish to use to build a surrogate model to increase accuracy in high posterior probability regions.</td>
</tr>
<tr>
<td><code>import_build_points_file</code></td>
<td>File containing points you wish to use to build a surrogate model.</td>
</tr>
<tr>
<td><code>posterior_adaptive</code></td>
<td>Adapt emulator model to increase accuracy in high posterior probability regions.</td>
</tr>
</tbody>
</table>

### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

#### 7.2.68.3.3.99 collocation_points

- Keywords Area
- `method`
- `bayes_calibration`
- `wasabi`
- `emulator`
- `pce`
- `orthogonal_least_interpolation`
- `collocation_points`

Number of collocation points used to estimate expansion coefficients

### Specification

**Alias:** none
**Argument(s):** INTEGER

### Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.

A corresponding sequence specification is documented at, e.g., `collocation_points_sequence` and `collocation_points_sequence`.
See Also

These keywords may also be of interest:

- collocation_points_sequence  method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence

7.2.68.3.3.100 tensor_grid

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none

Argument(s): INTEGERLIST

Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.3.3.101 reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.68.3.102 import_build_points_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td></td>
<td>custom_annotated</td>
<td>selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>selects freeform file format</td>
</tr>
</tbody>
</table>
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g., surrogate-based optimization, or implicitly, e.g., efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```plaintext
method
polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

Selects custom-annotated tabular file format

### Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format  

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The *annotated* format is the default for tabular export/import. To control which header row and columns are in the input/output, specify *custom.annotated*, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to *annotated* format, though *freeform* remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
    tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom.annotated header eval_id
```

Resulting tabular file:
7.2 method

7.2.68.3.3.104 header

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated

7.2.68.3.3.105 eval_id

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent *custom_annotated*

7.2.68.3.3.106  interface_id

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent *custom_annotated*

7.2.68.3.3.107  annotated

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   NO_ID   0.9  1.1  0.0002  0.26   0.76
2   NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3   NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2.68.3.3.108  freeform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to **annotated** format, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In **freeform**, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1 0.0001996404857  0.2601620081  0.759955
0.89991 1.1 0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.3.3.109  active_only

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
Adapt emulator model to increase accuracy in high posterior probability regions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

**Examples**

```plaintext
bayes_calibration queso
    chain_samples = 2000 seed = 348
    delayed_rejection
    emulator
        gaussian_process surfpack build_samples = 30
        posterior_adaptive max_iterations = 10
        proposal_covariance derivatives
```

**Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.**
7.2 method

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.3.3.113 normalized

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- normalized

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

Specification

**Alias**: none

**Argument(s)**: none

**Default**: PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.3.3.114 export_expansion_file

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

**Alias**: none

**Argument(s)**: STRING
Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consist only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.3.3.115 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none
Argument(s): none
Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.3.3.116 full_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- pce
- full_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none
Description

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.68.3.3.117  ml_pce

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce

Multilevel Polynomial Chaos Expansion as an emulator model.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multilevel expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
</tbody>
</table>
### Optional (Choose One) Basis Polynomial Family (Group 2)

<table>
<thead>
<tr>
<th>askey</th>
<th>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</th>
</tr>
</thead>
<tbody>
<tr>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Optional</th>
<th>normalized</th>
<th>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials. Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Optional (Choose One) Covariance Type (Group 3)</th>
<th>diagonal_covariance</th>
<th>Display only the diagonal terms of the covariance matrix.</th>
</tr>
</thead>
<tbody>
<tr>
<td>full_covariance</td>
<td>full_covariance</td>
<td>Display the full covariance matrix.</td>
</tr>
</tbody>
</table>

### Description

Selects a multilevel polynomial chaos expansion (ML PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using ML PCE as a surrogate within the Bayesian framework.

### See Also

These keywords may also be of interest:

- multilevel_polynomial_chaos

7.2.68.3.3.118 allocation_control

- Keywords Area
Sample allocation approach for multilevel expansions

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Multilevel Sample Allocation Control (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>estimator_variance</td>
<td>Variance of mean estimator within multilevel polynomial chaos</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rip_sampling</td>
<td>Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

**Description**

Multilevel expansions, including regression-based polynomial chaos expansion (PCE) and function train (FT) expansions, require a sample allocation strategy. Three options are currently available:

- allocation based on assuming a convergence rate for the estimator variance (for regression PCE)
- restricted isometry property (RIP) sampling (for regression PCE via compressed sensing)
- rank sampling (for FT)

**Default Behavior**

Current defaults, when allocation_control is not specified, are estimator variance for PCE and rank sampling for FT.

7.2.68.3.3.119 estimator_variance

- Keywords Area
  - method
  - bayes_calibration
  - wasabi
Variance of mean estimator within multilevel polynomial chaos

**Specification**

**Alias:** none  
**Argument(s):** none

| Child Keywords: |
|-----------------|-----------------|-----------------|
| Required/Optional | Description of Group | Dakota Keyword | Description |
| Optional          |                          | estimator_rate  | Rate of convergence of mean estimator variance within multilevel polynomial chaos |

**Description**

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
\text{Var} [\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
\text{Var} [\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}
\]

for free parameters \( \gamma \) and \( \kappa \), with default values that may be overridden as part of this specification block.

This approach is supported for regression-based PCE approaches (over-determined least squares, under-determined compressed sensing, or orthogonal least interpolation).

In practice, it can be challenging to estimate a smooth convergence rate for estimator variance in the presence of abrupt transitions in the quality of sparse recoveries. As a result, sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the convergence of the estimator variance, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.
Dakota Version 6.15 Reference Manual generated on November 5, 2021

method,
model_pointer = ‘HIERARCH’
multilevel_polynomial_chaos
  orthogonal_matching_pursuit
  expansion_order_sequence = 2
  pilot_samples = 10
  collocation_ratio = .9
  allocation_control
    estimator_variance estimator_rate = 2.5
  seed = 1237
  convergence_tolerance = .01

7.2.68.3.3.120  estimator_rate

  • Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • emulator
  • ml_pce
  • allocation_control
  • estimator_variance
  • estimator_rate

Rate of convergence of mean estimator within multilevel polynomial chaos

Specification

  Alias: none
  Argument(s): REAL
  Default: 2

Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

$$Var[\hat{Q}] = \frac{\sigma_Q^2}{N}$$

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

$$Var[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}$$

for free parameters $\gamma$ and $\kappa$.

The default values are $\gamma = 1$ and $\kappa = 2$ (adopts a more aggressive sample profile by assuming a faster convergence rate than Monte Carlo). This advanced specification option allows to user to specify $\kappa$, overriding the default.
Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Multilevel polynomial chaos with compressed sensing may allocate the number of samples per level based on the restricted isometry property (RIP), applied to recovery at level $l$:

$$N_l \geq s_l \log^3(s_l) L_l \log(C_l)$$

for sparsity $s$, cardinality $C$, and mutual coherence $L$. The adaptive algorithm starts from a pilot sample, shapes the profile based on observed sparsity, and iterates until convergence. In practice, RIP sampling levels are quite conservative, and a collocation ratio constraint ($N_l \leq rC_l$, where $r$ defaults to 2) must be enforced on the profile.

The algorithm relies on observed sparsity, it is appropriate for use with regularized solvers for compressed sensing. It employs orthogonal matching pursuit (OMP) by default and automatically activates cross-validation in order to choose the best noise parameter value for the recovery.

This capability is **experimental**. Sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the number of sparse coefficient sets recovered for each level, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```dakota
method, model_pointer = 'HIERARCH'
multilevel_polynomial_chaos orthogonal_matching_pursuit
  expansion_order_sequence = 2
  pilot_samples = 10
  collocation_ratio = .9
  allocation_control rip_sampling
  seed = 1237
  convergence_tolerance = .01
```
7.2.68.3.3.122 discrepancy_emulation

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- discrepancy_emulation

Formulation for emulation of model discrepancies.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** distinct

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Discrepancy Emulation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>recursive</td>
<td>distinct</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

**Description**

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.
Examples

method, multilevel_polynomial_chaos
  expansion_order_sequence = 2
  collocation_ratio = .9
  orthogonal_matching_pursuit
  discrepancy_emulation recursive

7.2.68.3.123  distinct

  • Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • emulator
  • ml_pce
  • discrepancy_emulation
  • distinct

Distinct formulation for emulation of model discrepancies.

Specification

Alias: paired

Argument(s): none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.3.124  recursive

  • Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • emulator
  • ml_pce
  • discrepancy_emulation
  • recursive

Recursive formulation for emulation of model discrepancies.
Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.3.3.125 expansion_order_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence

Sequence of expansion orders used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td></td>
</tr>
<tr>
<td>----------------------------------------------</td>
<td>-----------------------</td>
<td>----------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>basis_type</td>
<td>collocation_ratio</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
<td></td>
</tr>
<tr>
<td>expansion_samples_sequence</td>
<td></td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., expansion_samples.</td>
<td></td>
</tr>
</tbody>
</table>
### Keywords Area

| Optional | import_build_points_file | File containing points you wish to use to build a surrogate |

#### Description

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

#### See Also

These keywords may also be of interest:

- `expansion_order`

7.2.68.3.3.126  dimension_preference

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- `expansion_order_sequence`
- `dimension_preference`

A set of weights specifying the relative importance of each uncertain variable (dimension)

#### Specification

**Alias:** none

**Argument(s):** REALIST

**Default:** isotropic grids

#### Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.127 basis_type

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group PCE Basis Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max (i_1, \ldots, i_d) \leq p$$

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none
7.2 method

Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.68.3.3.130 adapted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>advancements</td>
<td></td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td>soft_convergence_limit</td>
<td></td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.3.3.131 advancements

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.132 soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.
7.2 method

Specification

**Alias:** none

**Argument(s):** INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.133  collocation_ratio

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **collocation_ratio**

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

**Alias:** none

**Argument(s):** REAL

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td><strong>orthogonal_matching_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>cross_validation</strong></td>
<td>Use cross validation to choose the ‘best’ polynomial order of a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>ratio_order</strong></td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the collocation_ratio allows for specification of a constant factor applied to \( N \) (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o \) for a polynomial chaos expansion). \[ 7.2.68.3.3.134 \]

<table>
<thead>
<tr>
<th>Optional</th>
<th>response_scaling</th>
<th>Perform bounds-scaling on response values prior to surrogate emulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
</tr>
</tbody>
</table>

### collocation_points_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
Specification

**Alias:** pilot_samples

**Argument(s):** INTEGERLIST

**Description**

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

**See Also**

These keywords may also be of interest:

- `collocation_points`

7.2.68.3.3.135 least_squares

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- `expansion_order_sequence`
- `collocation_ratio`
- `least_squares`

Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Optional (Choose One)

<table>
<thead>
<tr>
<th>LSQ Regression Approach (Group 1)</th>
<th>svd</th>
</tr>
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<tbody>
<tr>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
<td></td>
</tr>
<tr>
<td>equality_constrained</td>
<td></td>
</tr>
<tr>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
<td></td>
</tr>
</tbody>
</table>

### Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.68.3.3.136 svd

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_squares
- svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

### Specification

**Alias:** none

**Argument(s):** none

### Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

### orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp  
**Argument(s):** none

**Child Keywords:**
## 7.2.68.3.3.139 noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

### Specification

**Alias:** bp  
**Argument(s):** none

### Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.3.3.141 basis_pursuit_denoising

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

### Specification

**Alias:** bpdn  
**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
7.2 method

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.3.3.142 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.143 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
Specification

**Alias:** lars

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.3.3.144 noise_tolerance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **collocation_ratio**
- **least_angle_regression**
- **noise_tolerance**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
7.2 method 2551

7.2.68.3.3.145 least_absolute_shrinkage

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $\ell_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td></td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.3.3.146 noise_tolerance

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.147  \( l_2 \) penalty

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_absolute_shrinkage
- \( l_2 \) penalty

The \( l_2 \) penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The \( l_2 \) penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

*Alias*: none

*Argument(s):* none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.3.150 max_cv_order_candidates

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

Limit the number of cross-validation candidates for basis order
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.3.3.151 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^\omega$ where $c$ is the collocation_ratio and $\omega$ is the ratio_order.
Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

**7.2.68.3.3.152 response_scaling**
7.2 method

• expansion_order_sequence
• collocation_ratio
• use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.3.154 tensor_grid

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
7.2.68.3.3.155  reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.3.3.155  max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients
7.2 method

Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.3.3.157 expansion_samples_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
- expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the `expansion_samples_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_samples`

Specification

**Alias:** none  
**Argument(s):** INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

See Also

These keywords may also be of interest:

- `expansion_samples`
7.2.68.3.3.158  reuse_points

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
• expansion_order_sequence
• expansion_samples_sequence
• reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.3.3.159  import_build_points_file

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
• expansion_order_sequence
• import_build_points_file

File containing points you wish to use to build a surrogate
Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
<td></td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```plaintext
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.3.3.160  custom_annotated

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- expansion_order_sequence
Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

%eval_id  x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
... 
```

7.2.68.3.3.161  header

• Keywords Area

• method

• bayes_calibration

• wasabi

• emulator

• ml_pce

• expansion_order_sequence

• import_build_points_file

• custom_annotated

• header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

*Alias*: none  
*Argument(s)*: none  
*Default*: no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file
Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.3.3.164 annotated

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
• expansion_order_sequence
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id.

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...`

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.3.3.166  active_only

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• ml_pce
7.2 method

- expansion_order_sequence
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables. This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.3.167 orthogonal_least_interpolation

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

**Description**

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

7.2.68.3.3.168  collocation_points_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

**Specification**

**Alias**: none

**Argument(s)**: INTEGERLIST
7.2 method

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.68.3.3.169 tensor_grid

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none

Argument(s): INTEGERLIST

Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.3.3.170 reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

**7.2.68.3.171 import_build_points_file**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **ml_pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```dakota
method polynomial_chaos
   expansion_order = 4
   import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### Optional (Choose One)

<table>
<thead>
<tr>
<th>Tabular Format (Group 1)</th>
<th>custom_annotated</th>
<th>Selects custom-annotated tabular file format</th>
</tr>
</thead>
<tbody>
<tr>
<td>annotated</td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td>freeform</td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

| Optional                | active_only      | Import only active variables from tabular data file |

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.3.3.173 header

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.68.3.3.174 eval_id

- Keywords Area
- method
- bayes_calibration
- wasabi
Enable evaluation ID column in custom-annotated tabular file

**Specification**

Alias: none

Argument(s): none

Default: no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

Alias: none

Argument(s): none

Default: no interface_id column

**Description**

See description of parent `custom_annotated`
7.2.68.3.3.176  annotated

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface     x1    x2     obj_fn nln_ineq_con_1 nln_ineq_con_2
1      NO_ID       0.9    1.1    0.0002   0.26        0.76
2      NO_ID    0.90009 1.1  0.0001996404857 0.2601620081  0.759955
3      NO_ID    0.89991 1.1  0.0002003604863 0.2598380081  0.760045
...
```

7.2.68.3.3.177  freeform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Arguement(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
      freeform

Resulting tabular file:

<table>
<thead>
<tr>
<th>0.9</th>
<th>1.1</th>
<th>0.0002</th>
<th>0.26</th>
<th>0.76</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>
```

7.2.68.3.3.178  active_only

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.3.179 askey

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- polynomial_chaos  
- askey

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
Specification

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

**Description**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

7.2.68.3.3.182  **export_expansion_file**

- **Keywords Area**
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

**Specification**

**Alias:** none

**Argument(s):** STRING

**Description**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.3.3.183  **diagonal_covariance**

- **Keywords Area**
- method
- bayes_calibration
- wasabi
- emulator
- ml_pce
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.3.3.184 full_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mi_pce**
- **full_covariance**

Display the full covariance matrix

Specification

**Alias:** none

**Argument(s):** none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.

7.2.68.3.3.185 mf_pce

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**

Multifidelity Polynomial Chaos Expansion as an emulator model.
Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>quadrature_order_sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level_sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askkey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>
The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.

Display only the diagonal terms of the covariance matrix.

Display the full covariance matrix.

<table>
<thead>
<tr>
<th>Optional</th>
<th>Optional</th>
<th>Optional (Choose One)</th>
<th>Covariance Type (Group 3)</th>
<th>Export Expansion File</th>
<th>Full Covariance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>diagonal covariance</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Selects a multifidelity polynomial chaos expansion (MF PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF PCE as a surrogate within the Bayesian framework.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`

7.2.68.3.186  `p_refinement`

- `Keywords Area`
- `method`
- `bayes_calibration`
- `wasabi`
- `emulator`
- `mf_pce`
- `p_refinement`

Automatic polynomial order refinement

Specification

**Alias**: none

**Argument(s)**: none

**Default**: no refinement

**Child Keywords**: none
Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the max_iterations and convergence_tolerance method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the uniform refinement case, the quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.3.3.187 uniform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
<table>
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<th>Dakota Keyword</th>
<th>Description</th>
</tr>
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<tr>
<td>Required (Choose One)</td>
<td></td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>decay</td>
<td>Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.3.3.189 sobol

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.
Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized

**Description**

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

**Examples**

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.3.3.190 decay
  • Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • emulator
  • mf_pce
  • p_refinement
  • dimension_adaptive
  • decay
```

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive_p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```

**7.2.68.3.192** **max_refinement_iterations**

- **Keywords Area**
- **method**
- **bayes_calibration**

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive_p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```

**7.2.68.3.191** **generalized**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **p_refinement**
- **dimension_adaptive**
- **generalized**
Keywords Area

• wasabi
• emulator
• mf_pce
• max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

```plaintext
method,
polynomial_choos
dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

7.2.68.3.3.193 allocation_control

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_pce
• allocation_control

Sample allocation approach for multifidelity expansions

Specification

Alias: none
Argument(s): none
Child Keywords:
## Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

### Default Behavior

The default, when `allocation_control` is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

### 7.2.68.3.3.194 greedy

- **Keywords Area**
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - mf_pce
  - allocation_control
  - greedy

Sample allocation based on greedy refinement within multifidelity polynomial chaos

### Specification

**Alias:** none

**Argument(s):** none

### Description

Multifidelity polynomial chaos supports greedy refinement strategies, spanning regression and projection approaches for computing expansion coefficients. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within a unified competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any $z/p/\beta/\beta^*$ level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.
Examples

The following example of greedy multifidelity regression starts from a zeroth-order reference expansion (a constant) for each level, and generates candidate refinements for each level that are competed in an integrated greedy competition. The number of new samples for the incremented candidate expansion order is determined from the collocation ratio. In this case, the number of candidates for each level is limited to one uniform refinement of the current expansion order.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement uniform
    expansion_order_sequence = 0
    collocation_ratio = .9  seed = 160415
    orthogonal_matching_pursuit
    convergence_tolerance 1.e-2
```

The next example employs generalized sparse grids within a greedy multifidelity competition. Each modeling level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all model levels is competed within an integrated greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each model level and the number of candidates grows rapidly with random dimension and grid level.

```
method,
  model_pointer = 'HIERARCH'
  multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement dimension_adaptive generalized
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-8
```

7.2.68.3.3.195 discrepancy_emulation

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none
Default: distinct
Child Keywords:
In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.

### Examples

```plaintext
method,
    multilevel_polynomial_choas
    expansion_order_sequence = 2
    collocation_ratio = .9
    orthogonal_matching_pursuit
    discrepancy_emulation recursive
```

7.2.68.3.3.196  distinct

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **discrepancy_emulation**
- **distinct**

Distinct formulation for emulation of model discrepancies.
Specification

**Alias:** paired

**Argument(s):** none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.3.3.197  recursive

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.

Specification

**Alias:** none

**Argument(s):** none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples
7.2 method  

7.2.68.3.3.198 quadrature_order_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

Specification

Alias: none

 Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Each level entry of the quadrature_order_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., quadrature_order

See Also

These keywords may also be of interest:

- quadrature_order
A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

*Alias:* none

*Argument(s):* REALLIST

*Default:* isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay
7.2 method

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.3.201 non_nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none

Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.202 sparse_grid_level_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion
### Specification

**Alias:** none  
**Argument(s):** INTEGERLIST

#### Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>dimension_preference</td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
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</table>

#### Optional (Choose One)

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</thead>
<tbody>
<tr>
<td>Quadrature Rule Growth (Group 1)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td></td>
<td>unrestricted</td>
<td>Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.</td>
</tr>
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</table>

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

### See Also

These keywords may also be of interest:

- `sparse_grid_level`

7.2.68.3.3.203 `dimension_preference`

- **Keywords Area**
- **method**
A set of weights specifying the relative importance of each uncertain variable (dimension).

**Specification**

**Alias:** none

**Argument(s):** REALIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for sparse grid (\( i = 4l + 1 \)) or an order \( m \) for tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.3.205 unrestricted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

- **Alias**: none
- **Argument(s)**: none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for sparse grid (\( i = 4l + 1 \)) or an order \( m \) for tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none
Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

**7.2.68.3.3.208 expansion_order_sequence**

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence

Sequence of expansion orders used in a multi-stage expansion

### Specification

**Alias:** none  

**Argument(s):** INTEGERLIST

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<tr>
<th>Child Keywords:</th>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td>Optional</td>
<td>dimension_preference</td>
<td>basis_type</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension) Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td></td>
<td>Optional</td>
<td>collocation_ratio</td>
<td></td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td><strong>expansion_samples_sequence</strong></td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the <code>expansion_samples_sequence</code> applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., <code>expansion_samples</code></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**import_build_points_file**

File containing points you wish to use to build a surrogate

---

**Description**

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

**See Also**

These keywords may also be of interest:

- `expansion_order`

---

**dimension_preference**

A set of weights specifying the relative importance of each uncertain variable (dimension).

**Specification**

- **Alias:** none
- **Argument(s):** REALLIST
- **Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.210 basis_type

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>PCE Basis Type (Group 1)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none
Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.68.3.3.213 adapted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>advancements</td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>soft_convergence_limit</td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.3.3.214 advancements

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

Specification

Alias: none

Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.215 soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.
7.2 method

Specification

Alias: none
Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.3.3.216 collocation_ratio

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none
Argument(s): REAL

Child Keywords:

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<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
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<tr>
<td>Optional (Choose One)</td>
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<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
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<tr>
<td>Keyword</td>
<td>Description</td>
<td></td>
<td></td>
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<tr>
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<td>---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP).</td>
<td></td>
<td></td>
</tr>
<tr>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit ( \ell_1 )-minimization problem using linear programming.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising ( \ell_1 )-minimization problem using second order cone optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>least_angle_regression</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional cross_validation</td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional ratio_order</td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate $N$ from $n$ and $p$, the collocation_ratio allows for specification of a constant factor applied to $N$ (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with $N$ can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes $cN^o_{n+p}$).

### 7.2.68.3.3.217 collocation_points_sequence

- **Keywords Area**
- **method**
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
Specification

Alias: pilot_samples

Argument(s): INTEGERLIST

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.68.3.218 least_squares

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

Alias: none

Argument(s): none

Default: svd

Child Keywords:

<table>
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<th>Required/Optional Description of Group</th>
<th>Dakota Keyword</th>
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<td></td>
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</tbody>
</table>
7.2 method

<table>
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<tr>
<th>Optional (Choose One)</th>
<th>LSQ Regression Approach (Group 1)</th>
<th>svd</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>equality_constrained</td>
</tr>
</tbody>
</table>

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.68.3.3.219 svd

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_squares
- svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Specification

**Alias**: none

**Argument(s)**: none

Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none
Child Keywords:
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.3.3.222 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.223 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp  
**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.3.224  basis_pursuit_denoising

* Keywords Area
* method
* bayes_calibration
* wasabi
* emulator
* mf_pce
* expansion_order_sequence
* collocation_ratio
* basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn  
**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
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<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.3.3.225 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: $10^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.226 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
Specification

Alias: lars

Argument(s): none

Child Keywords:

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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.3.227 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_pce
• expansion_order_sequence
• collocation_ratio
• least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso

**Argument(s):** none

**Child Keywords:**

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<tr>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $\ell_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>l2_penalty</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_pce
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias**: none

**Argument(s)**: REALIST

**Default**: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.3.3.230  l2_penalty

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias**: none

**Argument(s)**: REAL

**Default**: 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>max_cv_order_-candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.3.3.232 noise_only

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.3.233 max_cv_order_candidates

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.3.3.234 ratio_order

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \(cN^o\) where \(c\) is the collocation_ratio and \(o\) is the ratio_order.
Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

---

7.2.68.3.3.236  use_derivatives

- Keywords Area  
- method  
- bayes_calibration  
- wasabi  
- emulator  
- mf_pce
Use derivative data to construct surrogate models

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.
Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.3.3.240 expansion_samples_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

Specification

Alias: none
Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
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<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

See Also

These keywords may also be of interest:

- expansion_samples
reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- expansion_samples_sequence
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

import_build_points_file

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- import_build_points_file

File containing points you wish to use to build a surrogate
Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
<table>
<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<td>Selects custom-annotated tabular file format</td>
</tr>
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<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```dakota
method polynomial_choas
   expansion_order = 4
   import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.3.3.243 custom_annotated

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
• import_build_points_file
  • custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

  • file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.3.3.244 header

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- expansion_order_sequence
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

### Specification

- **Alias**: none
- **Argument(s)**: none
- **Default**: no header

### Description

See description of parent `custom_annotated`
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Keywords Area

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

Description

See description of parent *custom_annotated*

7.2.68.3.3.247 annotated

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **expansion_order_sequence**
- **import_build_points_file**
- **annotated**

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- **file_formats**

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The *annotated* keyword can be used to explicitly specify this.

**Usage Tips**
• To specify pre-Dakota 6.1 tabular format, which did not include `interface_id`, specify `custom.annotated` header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though `freeform` remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

**Examples**

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

**7.2.68.3.3.248 freeform**

• Keywords Area

• method

• bayes_calibration

• wasabi

• emulator

• mf_pce

• expansion_order_sequence

• import_build_points_file

• freeform

Selects freeform file format

**Topics**

This keyword is related to the topics:

• file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9   1.1   0.0002   0.26   0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.3.3.249  active_only

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_pce
• expansion_order_sequence
• import_build_points_file
• active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.3.250 orthogonal_least_interpolation

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_pce
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:
## Keywords Area

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods expansion_order cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

7.2.68.3.251 collocation_points_sequence

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **orthogonal_least_interpolation**
- **collocation_points_sequence**

Sequence of collocation point counts used in a multi-stage expansion

### Specification

**Alias**: none

**Argument(s)**: INTEGERLIST
Description

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

See Also

These keywords may also be of interest:

- `collocation_points`

7.2.68.3.3.252 tensor_grid

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.3.3.253 reuse_points

- Keywords Area
- method
- bayes_calibration
- wasabi
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

### Specification

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

### Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

#### 7.2.68.3.254 import_build_points_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**

File containing points you wish to use to build a surrogate

### Specification

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

#### Default Behavior

By default, methods do not import points from a file.

#### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

#### Examples

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### Group 1

<table>
<thead>
<tr>
<th>Optional (Choose One)</th>
<th>Tabular Format</th>
<th>custom_annotated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

| Optional               | active_only                 | Import only active variables from tabular data file |

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.3.3.256  header

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.68.3.3.257  eval_id

- Keywords Area
- method
- bayes_calibration
- wasabi
Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no eval_id column

Description

See description of parent custom_annotated

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no interface_id column

Description

See description of parent custom_annotated
Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  NO_ID    0.9  1.1  0.0002  0.26  0.76
2  NO_ID    0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID    0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.3.3.260  freeform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
 0.9 1.1 0.0002 0.26 0.76
 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

**7.2.68.3.3.261 active_only**

- **Keywords Area**
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

**Topics**

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword \texttt{active\_only} indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.3.262 \texttt{askey}

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:
- \texttt{polynomial\_chaos}
- \texttt{wiener}
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- **polynomial_chaos**
- **askey**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
Specification

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

**Description**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.3.3.265 export_expansion_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **export_expansion_file**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

**Specification**

**Alias:** none

**Argument(s):** STRING

**Description**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.3.3.266 diagonal_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_pce**
- **diagonal_covariance**

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

Alias: none

Argument(s): none

Default: diagonal_c covariance for response vector > 10; else full covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.3.3.267 full_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_pce
- full_covariance

Display the full covariance matrix

Specification

Alias: none

Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full covariance is used to force Dakota to output the full covariance matrix.

7.2.68.3.3.268 sc

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc

Stochastic Collocation as an emulator model.
Specification

Alias: none
Argument(s): none
Child Keywords:
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<th>Description of Group</th>
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<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
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<td>h_refinement</td>
<td>Employ h-refinement to refine the grid</td>
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<td>Interpolation Grid Type (Group 2)</td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
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<td></td>
<td></td>
<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
</tr>
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<td></td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
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<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
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<tr>
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<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>
Use derivative data to construct surrogate models

Display only the diagonal terms of the covariance matrix

Display the full covariance matrix

**Description**

Selects stochastic collocation (SC) model to use in the Bayesian likelihood calculations. Most specification options are carried over for using SC as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- stoch_collocation

**Automatic polynomial order refinement**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**
7.2 method

| dimension_adaptive | Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’. |

**Description**

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional adaptive candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

**7.2.68.3.3.270 uniform**

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- `p_refinement`
- uniform

Refine an expansion uniformly in all dimensions.
Specification

**Alias:** none

**Argument(s):** none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

### 7.2.68.3.271 dimension_adaptive

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **sc**
- **p_refinement**
- **dimension_adaptive**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Dimension Adaptivity Estimation Approach (Group 1)</td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
</tr>
</tbody>
</table>
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

**7.2.68.3.3.272 sobol**

- **Keywords** Area 
- **method** 
- bayes_calibration 
- wasabi 
- emulator 
- sc 
- p_refinement 
- dimension_adaptive 
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** generalized

**Description**

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```

7.2.68.3.3.274  **h_refinement**

- **Keywords Area**
- **method**
- **bayes_calibration**

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Employ h-refinement to refine the grid

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>h-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
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<tr>
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<td></td>
<td><strong>dimension</strong>-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

**Description**

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_adaptive options.

**7.2.68.3.275 uniform**

- **Keywords Area**

- **method**
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**: | Required/Optional | Description of Group | Dakota Keyword | Dakota Keyword Description |
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<td>- bayes_calibration</td>
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<td>- wasabi</td>
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<td>- emulator</td>
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<td>- sc</td>
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<td>- h_refinement</td>
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<td>- uniform</td>
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</tr>
</tbody>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

**7.2.68.3.3.277 sobol**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **sc**
- **h_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```
method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p.refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.3.3.278 generalized

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- h.refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

**Alias**: none

**Argument(s)**: none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```
method,
  polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p.refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
7.2 method

7.2.68.3.279  local_adaptive

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- h_refinement
- local_adaptive

Planned future capability for local pointwise refinement within a generalized sparse grid.

Specification

**Alias:** none

**Argument(s):** none

Description

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

7.2.68.3.280  max_refinement_iterations

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

```plaintext
method,
polynomial_chaos
dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

7.2.68.3.3.281 quadrature_order

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
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<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order, and, optionally, dimension_preference). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a dimension_preference,
where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified \texttt{quadrature\_order} and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n}(p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the \texttt{quadrature\_order} specification supports an array input.

A corresponding sequence specification is documented at, e.g., \texttt{quadrature\_order\_sequence} and \texttt{quadrature\_order\_sequence}.

**See Also**

These keywords may also be of interest:

- \texttt{quadrature\_order\_sequence} and \texttt{method-multifidelity\_stoch\_collocation-quadrature\_order\_sequence}

7.2.68.3.3.282 \texttt{dimension\_preference}

- Keywords Area
- method
- bayes\_calibration
- wasabi
- emulator
- sc
- \texttt{quadrature\_order}
- \texttt{dimension\_preference}

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

\textbf{Alias:} none

\textbf{Argument(s):} REALLIST

\textbf{Default:} isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.3.283 nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- quadrature_order
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.3.284 non_nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules
7.2 method

Specification

**Alias:** none

**Argument(s):** none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.285 sparse_grid_level

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **sc**
- **sparse_grid_level**

Level to use in sparse grid integration or interpolation

Specification

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
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<td>Group 1</td>
<td>nodal</td>
<td>Employ a nodal sparse grid construction in stochastic collocation</td>
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## Keywords Area

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<th>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</th>
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<td>Enforce use of nested quadrature rules if available</td>
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<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit `non_nested` specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User’s Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User’s Manual for additional details). As for `quadrature_order`, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`.

### See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence

7.2.68.3.286 dimension_preference

- Keywords Area
- method
A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol  
- decay

**Employ a nodal sparse grid construction in stochastic collocation**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** nodal
Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.

7.2.68.3.3.288 hierarchical

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- sparse_grid_level
- hierarchical

Employ a hierarchical sparse grid construction

Specification

Alias: none
Argument(s): none

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.68.3.3.289 restricted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.
Specification

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)

**Description**

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2^l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.3.290 unrestricted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

**Alias:** none

**Argument(s):** none

**Description**

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2^l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences. To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.291 nested

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• sc
• sparse_grid_level
• nested

Enforce use of nested quadrature rules if available

Specification

Alias: none
Argument(s): none
Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.292 non_nested

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• sc
• sparse_grid_level
• non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none
7.2 method

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.293 piecewise

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- piecewise

Use piecewise local basis functions

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.3.294 askey

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.3.3.296  use_derivatives

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- use_derivatives

Use derivative data to construct surrogate models

Specification

**Alias**: none

**Argument(s)**: none

**Default**: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.3.3.297  diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. **diagonal_covariance** is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.3.3.298  full_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- sc
- full_covariance

Display the full covariance matrix

 Specification

**Alias:** none

**Argument(s):** none

Description

With a large number of responses, the covariance matrix can be very large. **full_covariance** is used to force Dakota to output the full covariance matrix.

7.2.68.3.3.299  mf_sc

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc

Multifidelity Stochastic Collocation as an emulator model.
7.2 method

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**: none
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<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description of Group</th>
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<tr>
<td>Optional (Choose One)</td>
<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
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<tr>
<td>Optional</td>
<td>h_refinement</td>
<td></td>
<td>Employ h-refinement to refine the grid</td>
</tr>
<tr>
<td>Optional</td>
<td>max_refinement_-iterations</td>
<td></td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td>allocation_control</td>
<td></td>
<td>Sample allocation approach for multifidelity expansions</td>
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<td>Optional</td>
<td>discrepancy_-_emulation</td>
<td></td>
<td>Formulation for emulation of model discrepancies</td>
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<tr>
<td>Required (Choose One)</td>
<td>Interpolation Grid Type (Group 2)</td>
<td>quadrature_order_-sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level_-sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables</td>
</tr>
</tbody>
</table>
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

<table>
<thead>
<tr>
<th>Optional</th>
<th>Covariance Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>wiener</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td></td>
<td>diagonal_c covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

**Description**

Selects a multifidelity stochastic collocation (MF SC) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF SC as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- `multifidelity_stoch_collocation`

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**
### Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.3.3.301 uniform

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
• mf_sc

• p_refinement

• uniform

Refine an expansion uniformly in all dimensions.

Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.3.3.302 dimension_adaptive

• Keywords Area

• method

• bayes_calibration

• wasabi

• emulator

• mf_sc

• p_refinement

• dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher 'importance' in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

#### 7.2.68.3.3.303 sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_sc**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

### Specification

- **Alias**: none
- **Argument(s)**: none
- **Default**: generalized

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Dimension Adapivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td><strong>sobol</strong></td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
<td><strong>generalized</strong></td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.3.3.304 generalized

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- p_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none
Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
7.2.68.3.3.305  h_refinement

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- h_refinement

Employ h-refinement to refine the grid

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional Required <em>(Choose One)</em></th>
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<th>Dakota Keyword</th>
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<tbody>
<tr>
<td></td>
<td>uniform</td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

**Description**

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_-adaptive options.
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords:**
## Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

### 7.2.68.3.3.308 sobol

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- h_refinement
- dimension_adaptive
- sobol

**Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.**

## Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.3.3.309  generalized

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- h_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none
Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
Planned future capability for local pointwise refinement within a generalized sparse grid.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

**7.2.68.3.3.310  local_adaptive**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_sc**
- **h_refinement**
- **local_adaptive**

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

method,  
  polynomial_chaos  
  dimension_adaptive generalized  
  p_refinement  
    max_refinement_iterations = 20  
    convergence_tol = 1.e-4  
    sparse_grid_level = 1

7.2.68.3.3.312 allocation_control

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- allocation_control

Sample allocation approach for multifidelity expansions

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Multifidelity Sample Allocation Control (Group 1)</td>
<td>greedy</td>
<td>Sample allocation based on greedy refinement within multifidelity stochastic collocation</td>
</tr>
</tbody>
</table>

Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

Default Behavior

The default, when allocation_control is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).
Sample allocation based on greedy refinement within multifidelity stochastic collocation

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Multifidelity stochastic collocation supports greedy refinement strategies using tensor and sparse grids for both nodal and hierarchical collocation approaches. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within an integrated competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.

**Examples**

The following example of greedy multifidelity stochastic collocation using nodal interpolation starts from a zeroth-order expansion (a constant) for each level, and generates uniform candidate refinements for each level that are competed in a greedy competition. The number of new samples for the incremented candidate expansion order is determined from the quadrature rules of the new sparse grid level. In this case, the number of candidates for each level is limited to one uniform refinement of the current sparse grid level.

```
method,  
  model_pointer = 'HIERARCH'  
  multifidelity_stoch_collocation  
  nodal  
  allocation_control greedy  
  p_refinement uniform  
  sparse_grid_level_sequence = 0 unrestricted  
  convergence_tolerance 1.e-3
```

The next example employs generalized sparse grids and hierarchical interpolation. Each level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all levels is competed within a unified greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each level and the number of candidates grows rapidly with random dimension and grid level.
method,
    model_pointer = 'HIERARCH'
    multifidelity_stoch_collocation
    hierarchical
    allocation_control greedy
    p_refinement dimension_adaptive generalized
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-8

7.2.68.3.3.314 discrepancy_emulation

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_sc
• discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none
Default: distinct
Child Keywords:

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<td>distinct</td>
<td>Distinct formulation for emulation of model discrepancies.</td>
</tr>
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</table>

| | recursive | Recursive formulation for emulation of model discrepancies. |

Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. distinct emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. recursive emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.
The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

Examples

```
method,
    multilevel_polynomial_chaos
    expansion_order_sequence = 2
    collocation_ratio = .9
    orthogonal_matching_pursuit
    discrepancy_emulation recursive
```

Distinct formulation for emulation of model discrepancies.

Specification

**Alias:** paired

**Argument(s):** none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

```
7.2.68.3.3.315    recursive
    Keywords Area
    method
    bayes_calibration
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Recursive formulation for emulation of model discrepancies.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

**Examples**

7.2.68.3.3.317  quadrature_order_sequence

- **Keywords Area**
  - method
  - bayes_calibration
  - wasabi
  - emulator
  - mf_sc
  - quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

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Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Optional (Choose One)

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<tbody>
<tr>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Each level entry of the `quadrature_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `quadrature_order`

### See Also

These keywords may also be of interest:

- `quadrature_order`

### Specification

- **Alias:** none
- **Argument(s):** REALLIST
- **Default:** isotropic grids
7.2 method

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.3.3.319 nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available

Specification

Alias: none

Argument(s): none

Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.3.3.320 non_nested

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules
Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.321 sparse_grid_level_sequence

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST

Child Keywords:

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### 7.2 method

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<td></td>
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<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

### See Also

These keywords may also be of interest:

- `sparse_grid_level`

### 7.2.68.3.3.322 dimension_preference

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- `sparse_grid_level_sequence`
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- `sobol`
- `decay`

7.2.68.3.3.323 nodal

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_sc**
- **sparse_grid_level_sequence**
- **nodal**

Employ a nodal sparse grid construction in stochastic collocation

Specification

**Alias:** none

**Argument(s):** none

**Default:** nodal

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.
Employ a hierarchical sparse grid construction

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g., mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.3.3.326 unrestricted

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none
Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none  
**Argument(s):** none
Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.3.3.329  piecewise

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_sc
• piecewise

Use piecewise local basis functions

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.3.3.330 askey

• Keywords Area
• method
• bayes_calibration
• wasabi
• emulator
• mf_sc
• askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

7.2.68.3.3.331 wiener

- **Keywords Area**
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.3.3.332 use_derivatives

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.3.3.333 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- wasabi
- emulator
- mf_sc
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.3.3.34 full_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **emulator**
- **mf_sc**
- **full_covariance**

Display the full covariance matrix

Specification

**Alias:** none

**Argument(s):** none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.

7.2.68.3.4 standardized_space

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **standardized_space**

Perform Bayesian inference in standardized probability space

Specification

**Alias:** none

**Argument(s):** none
Description

This option transforms the inference process (MCMC sampling and any emulator model management) into a standardized probability space.

The variable transformations performed are as described in askey.

Default Behavior

The default for the Gaussian process and no emulator options is to perform inference in the original probability space (no transformation). Polynomial chaos and stochastic collocation emulators, on the other hand, are always formed in standardized probability space, such that the inference process is also performed in this standardized space.

Expected Output

The user will see the truth model evaluations performed in the original space, whereas any method diagnostics relating to the MCMC samples (e.g., QUESO data in the outputData directory) will report points and response data (response gradients and Hessians, if present, will differ but response values will not) that correspond to the transformed space.

Usage Tips

Selecting standardized_space generally has the effect of scaling the random variables to be of comparable magnitude, which can improve the efficiency of the Bayesian inference process.

Examples

```plaintext
method,
    bayes_calibration queso
    samples = 2000 seed = 348
    dram
    standardized_space
```

7.2.68.3.5 data_distribution

- Keywords Area
- method
- bayes_calibration
- wasabi
- data_distribution

(Experimental Capability) Specify the distribution of the experimental data

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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7.2.68.3.5.1  gaussian

- Keywords Area
- method
- bayes_calibration
- wasabi
- data_distribution
- gaussian

(Experimental Capability) Gaussian error distribution

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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7.2.68.3.5.2  means

- Keywords Area
- method
- bayes_calibration
- wasabi
- data_distribution
- gaussian
- means

(Experimental Capability) Means of Gaussian error distribution

**Specification**

**Alias:** none

**Argument(s):** REALLIST
7.2.68.3.5.3 covariance

• Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • data_distribution
  • gaussian
  • covariance

(Experimental Capability) Covariance of a Gaussian error distribution

Specification

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

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<td>(Experimental Capability) Symmetric positive definite error covariance</td>
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7.2.68.3.5.4 diagonal

• Keywords Area
  • method
  • bayes_calibration
  • wasabi
  • data_distribution
  • gaussian
  • covariance
  • diagonal

(Experimental Capability) Diagonal error covariance

Specification

**Alias:** none

**Argument(s):** none
7.2.68.3.5.5 matrix

- Keywords Area
- method
- bayes_calibration
- wasabi
- data_distribution
- gaussian
- covariance
- matrix

(Experimental Capability) Symmetric positive definite error covariance

Specification

Alias: none
Argument(s): none

7.2.68.3.5.6 obs_data_filename

- Keywords Area
- method
- bayes_calibration
- wasabi
- data_distribution
- obs_data_filename

(Experimental Capability) Filename from which to read experimental data

Specification

Alias: none
Argument(s): STRING

7.2.68.3.6 posterior_samples_import_filename

- Keywords Area
- method
- bayes_calibration
- wasabi
- posterior_samples_import_filename

(Experimental Capability) Filename for samples at which the user would like the posterior density calculated
**Specification**

**Alias:** none

**Argument(s):** STRING

### 7.2.68.3.7  generate_posterior_samples

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **generate_posterior_samples**

(Experimental Capability) Generate random samples from the posterior density

Description: This keyword will result in samples from the prior that have a high posterior density being printed to a file called 'psamples.txt' as a default, or to a file specified by the `posterior_samples_export_filename`.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
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<td>(Experimental Capability) Filename for the exported posterior samples</td>
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### 7.2.68.3.7.1  posterior_samples_export_filename

- **Keywords Area**
- **method**
- **bayes_calibration**
- **wasabi**
- **generate_posterior_samples**
- **posterior_samples_export_filename**

(Experimental Capability) Filename for the exported posterior samples

**Specification**

**Alias:** none

**Argument(s):** STRING
7.2.68.3.8 evaluate_posterior_density

- Keywords Area
- method
- bayes_calibration
- wasabi
- evaluate_posterior_density

(Experimental Capability) Evaluate the posterior density and output to the specified file

**Description:** This keyword will allow the evaluation of the posterior density for all of the prior samples, typically specified with the number given in `pushforward_samples`. The density will be printed either to a file named `pdens.txt` as a default, or to the file specified in `posterior_density_export_filename`.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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7.2.68.3.8.1 posterior_density_export_filename

- Keywords Area
- method
- bayes_calibration
- wasabi
- evaluate_posterior_density
- posterior_density_export_filename

(Experimental Capability) Filename for the exported posterior density

**Specification**

**Alias:** none

**Argument(s):** STRING

7.2.68.4 dream

- Keywords Area
- method
DREAM (Differential Evolution Adaptive Metropolis)

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Child Keywords:

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<td>Optional</td>
<td></td>
<td>gr_threshold</td>
<td>Convergence tolerance for the Gelman-Rubin statistic</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>jump_step</td>
<td>Number of generations a long jump step is taken</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>emulator</td>
<td>Use an emulator or surrogate model to evaluate the likelihood function</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>standardized_space</td>
<td>Perform Bayesian inference in standardized probability space</td>
</tr>
</tbody>
</table>
### 7.2 method

<table>
<thead>
<tr>
<th>Optional</th>
<th>export_chain_points_file</th>
<th>Export the MCMC chain to the specified filename</th>
</tr>
</thead>
</table>

#### Description

The DiffeRential Evolution Adaptive Metropolis algorithm is described in[89]. For the DREAM method, one can define the number of chains used with chains (minimum 3). The total number of generations per chain in DREAM is the number of samples (samples) divided by the number of chains (chains). The number of chains randomly selected to be used in the crossover each time a crossover occurs is crossover_chain_pairs. There is an extra adaptation during burn-in, in which DREAM estimates a distribution of crossover probabilities that favors large jumps over smaller ones in each of the chains. Normalization is required to ensure that all of the input dimensions contribute equally. In this process, a discrete number of candidate points for each crossover value is generated. This parameter is num_cr. The gr_threshold is the convergence tolerance for the Gelman-Rubin statistic which will govern the convergence of the multiple chain process. The integer jump_step forces a long jump every jump_step generations. For more details about these parameters, see[89].

**Attention:** While the emulator specification for DREAM includes the keyword posterior_adaptive, it is not yet operational.

#### 7.2.68.4.1 chain_samples

- **Keywords** Area, method, bayes_calibration, dream, chain_samples

Number of Markov Chain Monte Carlo posterior samples

**Specification**

- **Alias:** samples
- **Argument(s):** INTEGER
- **Default:** method-dependent

**Description**

The chain_samples keyword indicates the number of draws from the posterior distribution to perform. When an emulator is active, this will be the number of samples on the constructed surrogate model.

**Default Behavior**

The default number of chain samples is method-dependent. QUESO and MUQ methods use 1000. DREAM uses (number of generations) x (number of chains), resulting in chain_samples close to that specified.

**Usage Tips**

MCMC methods typically require a large number of chain samples to converge, often thousands to millions.

**Examples**

```
method
    bayes_calibration queso
    chain_samples = 20000
```
7.2.68.4.2 seed

• Keywords Area
• method
• bayes_calibration
• dream
• seed

Seed of the random number generator

Specification

Alias: none

Argument(s): INTEGER

Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.68.4.3 chains

• Keywords Area
• method
• bayes_calibration
• dream
• chains

Number of chains in DREAM
7.2.68.4.4 num_cr

Number of candidate points for each crossover.

Topics
This keyword is related to the topics:

- bayesian_calibration
7.2.68.4.5 crossover_chain_pairs

- Keywords Area
- method
- bayes_calibration
- dream
- crossover_chain_pairs

Number of chains used in crossover.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): INTEGER

Default: 3

Description

The number of chains randomly selected to be used in the crossover each time a crossover occurs.

7.2.68.4.6 gr_threshold

- Keywords Area
- method
- bayes_calibration
- dream
- gr_threshold

Convergence tolerance for the Gelman-Rubin statistic

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): REAL

Default: 1.2
7.2 method

Description

The gr_threshold is the convergence tolerance for the Gelman-Rubin statistic which will govern the convergence of the multiple chain process.

7.2.68.4.7 jump_step

- Keywords Area
- method
- bayes_calibration
- dream
- jump_step

Number of generations a long jump step is taken

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none
Argument(s): INTEGER
Default: 5

Description

The integer jump_step forces a long jump every jump_step generations in DREAM.

7.2.68.4.8 emulator

- Keywords Area
- method
- bayes_calibration
- dream
- emulator

Use an emulator or surrogate model to evaluate the likelihood function

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

This keyword describes the type of emulator used when calculating the likelihood function for the Bayesian calibration. The emulator can be a Gaussian process, polynomial chaos expansion, or stochastic collocation.

#### 7.2.68.4.8.1 gaussian_process

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **gaussian_process**

Gaussian Process surrogate model

### Specification

- **Alias:** kriging
- **Argument(s):** none
- **Default:** Surfpack Gaussian process
- **Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Emulator Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
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<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
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<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
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<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
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<td>Description</td>
</tr>
<tr>
<td>Required/Optional (Choose One)</td>
<td>Emulator Type (Group 1)</td>
<td>Dakota Keyword</td>
<td>Description</td>
</tr>
</tbody>
</table>
Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the `surfpack` keyword.

An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

### 7.2.68.4.8.2 surfpack

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **gaussian_process**
- **surfpack**

Use the Surfpack version of Gaussian Process surrogates

### Specification

**Alias:** none

**Argument(s):** none
Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See `optimization_method`.
   The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See `trend`.

3. Correlation Lengths:
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See `correlation_lengths`.

4. Ill-conditioning
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using

5. Gradient Enhanced Kriging (GEK).
   The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   See notes in the Theory section.

Theory

Gradient Enhanced Kriging

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.
Select the built in Gaussian Process surrogate

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

A second version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the **dakota** version is deprecated and intended to be removed in a future release.**

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota's. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the **gaussian_process** **dakota** model can be found in [60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a **point_selection** option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the **point_selection** option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.
Specification

Alias: none
Argument(s): INTEGER

Description

The number of build points or training points used in the initial phase of an algorithm or model construction. Typically these are the initial set of data points used to construct (train) a surrogate model (emulator). If the number of build_samples is less than the minimum number of points required to build the surrogate, Dakota will augment the samples to obtain the minimum required.

Examples

Perform GP-based adaptive importance sampling, building the GP with 100 points and then performing 100 approximate evaluations to evaluate the probability.

```
method
  gpais
    build_samples = 100
    samples_on_emulator = 100
    max_iterations = 5
    response_levels = -1.065
```

7.2.68.4.8.5  posterior_adaptive

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- gaussian_process
- posterior_adaptive

Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none

Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.
7.2 method

Examples

bayes_calibration queso
  chain_samples = 2000 seed = 348
  delayed_rejection
  emulator
    gaussian_process surfpack build_samples = 30
    posterior_adaptive max_iterations = 10
    proposal_covariance derivatives

7.2.68.4.8.6 import_build_points_file

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- gaussian_process
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file

Argument(s): STRING

Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

The import_build_points_file allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.
Default Behavior
By default, methods do not import points from a file.

Usage Tips
Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples
```
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.4.8.7 custom_annotated

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- gaussian_process
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics
This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>%eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.68.4.8.8 header

• Keywords Area
  • method
Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated
7.2 method

7.2.68.4.8.10  interface_id

  • Keywords Area
  • method
  • bayes_calibration
  • dream
  • emulator
  • gaussian_process
  • import_build_points_file
  • custom_annotated
  • interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Aliased: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.4.8.11  annotated

  • Keywords Area
  • method
  • bayes_calibration
  • dream
  • emulator
  • gaussian_process
  • import_build_points_file
  • annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

  • file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include `interface_id`, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to *annotated* format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading `eval_id` and `interface_id` columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.68.4.8.12 freeform

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
• **gaussian_process**
• **import_build_points_file**
• **freeform**

Selects freeform file format

**Topics**

This keyword is related to the topics:

• **file_formats**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### 7.2.68.4.8.13 active_only

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **gaussian_process**
- **import_build_points_file**
- **active_only**

Import only active variables from tabular data file

#### Topics

This keyword is related to the topics:

- **file_formats**

#### Specification

**Alias:** none  
**Argument(s):** none

#### Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

### 7.2.68.4.8.14 pce

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**

Polynomial Chaos Expansion surrogate model
7.2 method

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>max_refinement_-iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Group 1</td>
<td>quadratic_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cubature_integrand</td>
<td>Cubature using Stroud rules and their extensions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_order</td>
<td>The (initial) order of a polynomial expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_-interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td><strong>Optional (Choose One)</strong></td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
</tbody>
</table>
### Description

Selects a polynomial chaos expansion (PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using PCE as a surrogate within the Bayesian framework.

### See Also

These keywords may also be of interest:

- polynomial_chaos

---

**Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.**

#### Optional

**normalized**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

**export_expansion_file**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.

#### Optional (Choose One)

**Covariance Type (Group 3)**

- **diagonal_covariance**
  - Display only the diagonal terms of the covariance matrix
- **full_covariance**
  - Display the full covariance matrix

---

**Automatic polynomial order refinement**
**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no refinement  

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>p-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.</td>
</tr>
</tbody>
</table>

**Description**

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the max_iterations and convergence_tolerance method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the uniform refinement case, the quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.
7.2.68.4.8.16  uniform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.

Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.4.8.17  dimension_adaptive

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

Specification

Alias: none
Argument(s): none
Child Keywords:
## Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

### sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.
7.2 method

Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4

7.2.68.4.8.19 decay

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

Specification

Alias: none
Argument(s): none

Description

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

**Examples**

```
method,  
polynomial_chaos  
sparse_grid_level = 3  
dimension_adaptive p_refinement generalized  
max_iterations = 20  
convergence_tol = 1.e-4
```

7.2.68.4.8.21  **max_refinement_iterations**

- Keywords Area
- method
- bayes_calibration
7.2 method

- dream
- emulator
- pce
- max_refinement_iterations

Maximum number of expansion refinement iterations

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

Limits the number of times the expansion can be refined under various refinement strategies.

Examples

```
method, polynomial_chaos
dimension_adaptive generalized p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

7.2.68.4.8.22 quadrature_order

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- quadrature_order

Order for tensor-products of Gaussian quadrature rules

Specification

Alias: none
Argument(s): INTEGER
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>dimension_preference</td>
<td></td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>nested</td>
<td></td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>non_nested</td>
<td></td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

**Description**

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with `quadrature_order`, and, optionally, `dimension_preference`). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of `p_refinement` or the case of an explicit `nested` override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a `dimension_preference`, where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified `quadrature_order` and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order $p_i$ of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision $2m_i - 1$, $p_i$ is one less than the quadrature order $m_i$ in each dimension (a one-dimensional expansion contains the same number of terms, $p_i + 1$, as the number of Gauss points). The total number of terms, $N$, in a tensor-product expansion involving $n$ uncertain input variables is

$$N = 1 + P = \prod_{i=1}^{n}(p_i + 1)$$

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the `quadrature_order` specification supports an array input.

A corresponding sequence specification is documented at, e.g., `quadrature_order_sequence` and `quadrature_order_sequence`.

**See Also**

These keywords may also be of interest:

- `quadrature_order_sequence` and `method-multifidelity_stoch_collocation-quadrature_order_sequence`

7.2.68.4.8.23  `dimension_preference`

- Keywords Area
- method
- bayes_calibration
- dream
A set of weights specifying the relative importance of each uncertain variable (dimension)

 Specification

 Alias: none
 Argument(s): REALLIST
 Default: isotropic grids

 Description

 A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

 See Also

 These keywords may also be of interest:

 - sobol
 - decay

7.2.68.4.8.24 nested

 Specification

 Alias: none
 Argument(s): none
 Default: quadrature: non_nested unless automated refinement; sparse grids: nested
Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.25 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules

Specification

**Alias:** none

**Argument(s):** none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.4.8.26 sparse_grid_level

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level

Level to use in sparse grid integration or interpolation

Specification

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**
## Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit `non_nested` specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User’s Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User's Manual for additional details). As for quadrature_order, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`.

### See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence
7.2.68.4.8.27  dimension_preference

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

Specification

Alias: none

Argument(s): REALLIST

Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification lead to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.28  restricted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.
7.2 method

Specification

Alias: none
Argument(s): none
Default: restricted (except for generalized sparse grids)

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.29 unrestricted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none
Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences. To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.30 nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level
- nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.31 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- sparse_grid_level
- non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none
7.2 method

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.4.8.32 cubature_integrand

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- cubature_integrand

Cubature using Stroud rules and their extensions

Specification

Alias: none

Argument(s): INTEGER

Description

Multi-dimensional integration by Stroud cubature rules [80] and extensions [94], as specified with cubature_integrand. A total-order expansion is used, where the isotropic order \( p \) of the expansion is half of the integrand order, rounded down. The total number of terms \( N \) for an isotropic total-order expansion of order \( p \) over \( n \) variables is given by

\[
N = 1 + P = 1 + \sum_{s=1}^{p} \frac{1}{s!} \prod_{r=0}^{s-1} (n+r) = \frac{(n+p)!}{n!p!}
\]

Since the maximum integrand order is currently five for normal and uniform and two for all other types, at most second- and first-order expansions, respectively, will be used. As a result, cubature is primarily useful for global sensitivity analysis, where the Sobol’ indices will provide main effects and, at most, two-way interactions. In addition, the random variable set must be independent and identically distributed (iid), so the use of askey or wiener transformations may be required to create iid variable sets in the transformed space (as well as to allow usage of the higher order cubature rules for normal and uniform). Note that global sensitivity analysis often assumes uniform bounded regions, rather than precise probability distributions, so the iid restriction would not be problematic in that case.

7.2.68.4.8.33 expansion_order

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order

The (initial) order of a polynomial expansion
Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimensional_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td></td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>collocation_points</td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td></td>
<td></td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_samples</td>
<td>Number of simulation samples used to estimate the expected value of a set of PCE coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>posterior_adaptive</td>
<td>Adapt emulator model to increase accuracy in high posterior probability regions</td>
</tr>
</tbody>
</table>

Description

When the expansion_order for a polynomial chaos expansion is specified, the coefficients may be computed by integration based on random samples or by regression using either random or sub-sampled tensor product.
quadrature points.

Multidimensional integration by Latin hypercube sampling (specified with `expansion_samples`). In this case, the expansion order \( p \) cannot be inferred from the numerical integration specification and it is necessary to provide an `expansion_order` to specify \( p \) for a total-order expansion.

Linear regression (specified with either `collocation_points` or `collocation_ratio`). A total-order expansion is used and must be specified using `expansion_order` as described in the previous option. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the `collocation_ratio` allows for specification of a constant factor applied to \( N \) (e.g., `collocation_ratio = 2` produces samples = \( 2N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using `ratio_order`. In this case, the number of samples becomes \( cN^o \) where \( c \) is the `collocation_ratio` and \( o \) is the `ratio_order`. The `use_derivatives` flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o \)). When admissible, a constrained least squares approach is employed in which response values are first reproduced exactly and error in reproducing response derivatives is minimized. Two collocation grid options are supported: the default is Latin hypercube sampling ("point collocation"), and an alternate approach of "probabilistic collocation" is also available through inclusion of the `tensor_grid` keyword. In this alternate case, the collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

If `collocation_points` or `collocation_ratio` is specified, the PCE coefficients will be determined by regression. If no regression specification is provided, appropriate defaults are defined. Specifically SVD-based least-squares will be used for solving over-determined systems and under-determined systems will be solved using LASSO. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares. Technical information on the various methods listed below can be found in the Linear regression section of the Theory Manual. Some of the regression methods (OMP, LASSO, and LARS) are able to produce a set of possible PCE coefficient vectors (see the Linear regression section in the Theory Manual). If cross validation is inactive, then only one solution, consistent with the `noise_tolerance`, will be returned. If cross validation is active, Dakota will choose between possible coefficient vectors found internally by the regression method across the set of expansion orders (1,...,`expansion_order` and the set of specified noise tolerances and return the one with the lowest cross validation error indicator.

### 7.2.68.4.8.34 dimension_preference

- **Keywords** Area
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **expansion_order**
- **dimension_preference**

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids
Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.35 basis_type

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group PCE Basis Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.

7.2.68.4.8.36 tensor_product

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- basis_type
- tensor_product

Use a tensor-product index set to construct a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none

Description

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max (i_1, \ldots, i_d) \leq p$$

7.2.68.4.8.37 total_order

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.68.4.8.38 adapted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
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<th>Description</th>
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</thead>
<tbody>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>advancements</td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>soft_convergence_limit</td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

**7.2.68.4.8.39 advancements**

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.4.8.40  soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.

Specification

Alias: none

Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.4.8.41  collocation_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points

Number of collocation points used to estimate expansion coefficients

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:
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<th>Dakota Keyword</th>
<th>Description</th>
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<td>7.2 method</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
</tr>
<tr>
<td></td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit (\ell_1)-minimization problem using linear programming.</td>
</tr>
<tr>
<td></td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising (\ell_1)-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td>least_angle_regression</td>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td></td>
<td>least_absolute_shrinkage</td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
<tr>
<td>Optional</td>
<td>cross_validation</td>
<td>Use cross validation to choose the ‘best’ polynomial order of a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>ratio_order</td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
</tr>
<tr>
<td>Optional</td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation.</td>
</tr>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models.</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients.</td>
</tr>
</tbody>
</table>

**Description**

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.

A corresponding sequence specification is documented at, e.g., `collocation_points_sequence` and `collocation_points_sequence`.

**See Also**

These keywords may also be of interest:

- `collocation_points_sequence` method-multilevel_polynomial_chaos-expansion_order_sequence-collocation-
Compute the coefficients of a polynomial expansion using least squares

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** svd

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<td>equality constrained</td>
<td>svd</td>
<td>Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.
Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Specification

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.
7.2 method

Specification

Alias: none
Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.4.8.45 orthogonal_matching_pursuit

- Keywords Area
- method
- bayes calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

Specification

Alias: omp
Argument(s): none

Child Keywords:

<table>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.4.8.46 noise_tolerance

- Keywords Area
- method
- bayes_calibration
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias**: none

**Argument(s)**: REALLIST

**Default**: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.47 basis_pursuit

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_points**
- **basis_pursuit**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias**: bp

**Argument(s)**: none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
7.2.68.4.8.48 basis_pursuit_denoising

• Keywords Area
  • method
  • bayes_calibration
  • dream
  • emulator
  • pce
  • expansion_order
  • collocation_points
  • basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

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<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.4.8.49 noise_tolerance

• Keywords Area
  • method
  • bayes_calibration
  • dream
  • emulator
  • pce
  • expansion_order
  • collocation_points
  • basis_pursuit_denoising
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.50  least_angle_regression

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_points**
- **least_angle_regression**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

**Specification**

**Alias:** lars  
**Argument(s):** none  

**Child Keywords:**

<table>
<thead>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
7.2.68.4.8.51 noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.52 least_absolute_shrinkage

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso

**Argument(s):** none

**Child Keywords:**

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

**7.2.68.4.8.53 noise_tolerance**

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points
- least_absolute_shrinkage
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### Specification

**Alias:** none

**Argument(s):** REALIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

### Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias**: none

**Argument(s)**: REAL

**Default**: 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.

Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:
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<tr>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_-candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

7.2.68.4.8.56 noise_only

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- noise_only

Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.
7.2 method

7.2.68.4.8.57 max_cv_order_candidates

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none

Argument(s): INTEGER

Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train
Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

**Specification**

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

**Description**

When using regression type methods (specified with either `collocation_points` or `collocation_ratio`), a total-order expansion can be specified using `expansion_order`. To avoid requiring the user to calculate $N$ from $n$ and $p$), the `collocation_ratio` allows for specification of a constant factor applied to $N$ (e.g., `collocation_ratio = 2`) produces samples = $2N$). In addition, the default linear relationship with $N$ can be overridden using a real-valued exponent specified using `ratio_order`. In this case, the number of samples becomes $cN^{\circ}$ where $c$ is the `collocation_ratio` and $\circ$ is the `ratio_order`.

Perform bounds-scaling on response values prior to surrogate emulation.
7.2 method

Specification

**Alias:** none
**Argument(s):** none
**Default:** use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.68.4.8.60 use_derivatives

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **expansion_order**
- **collocation_points**
- **use_derivatives**

Use derivative data to construct surrogate models

Specification

**Alias:** none
**Argument(s):** none
**Default:** use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.
Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
7.2 method

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points` file) can be reused.

7.2.68.4.8.63 max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_points
- max_solver_iterations

Maximum iterations in determining polynomial coefficients

Specification

Alias: none

Argument(s): INTEGER

Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.4.8.64 collocation_ratio

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none

Argument(s): REAL

Child Keywords:

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<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
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<td></td>
<td>orthogonal_matching_pursuit</td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
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<tr>
<td></td>
<td></td>
<td>basis_pursuit</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
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<td></td>
<td></td>
<td>basis_pursuit_denoising</td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td>Method</td>
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<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td></td>
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</tr>
<tr>
<td><code>least_angle_regression</code></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><code>least_absolute_shrinkage</code></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
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<tr>
<td><code>cross_validation</code></td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
<td></td>
<td></td>
</tr>
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<td><code>ratio_order</code></td>
<td>Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
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<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
<td></td>
<td></td>
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</table>
Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2, produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( \frac{cN^o}{p+1} \)).

7.2.68.4.8.65 least_squares

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

**Alias:** none

**Argument(s):** none

**Default:** svd

**Child Keywords:**

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<tr>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Optional/Choose One</td>
<td>Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.</td>
<td>svd</td>
<td></td>
</tr>
</tbody>
</table>
7.2 method

| equality_constrained | Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares. |

Description

Compute the coefficients of a polynomial expansion using least squares. Specifically SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

7.2.68.4.8.66 svd

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• pce
• expansion_order
• collocation_ratio
• least_squares
• svd

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition.

Specification

Alias: none

Argument(s): none

Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.

7.2.68.4.8.67 equality_constrained

• Keywords Area
• method
• bayes_calibration
• dream
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

7.2.68.4.8.68  *orthogonal_matching_pursuit*

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
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<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.4.8.69 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.70 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.
Specification

Alias: bp
Argument(s): none

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.4.8.71 basis_pursuit_denoising

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

Specification

Alias: bpdn
Argument(s): none

Child Keywords:

<table>
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<th>Dakota Keyword Description</th>
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<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.
7.2.68.4.8.72  noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.73  least_angle_regression

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

Specification

Alias: lars

Argument(s): none

Child Keywords:
### Keywords Area

<table>
<thead>
<tr>
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<th>Dakota Keyword Description</th>
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<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.4.8.74 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.75 least_absolute_shrinkage

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
Compute the coefficients of a polynomial expansion by using the LASSO problem.

Specification

Alias: lasso
Argument(s): none

Child Keywords:

<table>
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</thead>
<tbody>
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<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
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<tr>
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<td></td>
<td>l2_penalty</td>
<td>The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the LASSO problem.

7.2.68.4.8.76 noise_tolerance

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• pce
• expansion_order
• collocation_ratio
• least_absolute_shrinkage
• noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### 7.2.68.4.8.77 l2_penalty

- **Keywords Area**
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- least_absolute_shrinkage
- l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.

### 7.2.68.4.8.78 cross_validation

- **Keywords Area**
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Child Keywords**:

<table>
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<tr>
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<td>noise_only</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_-candidates</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.

- **noise_only**
  - Keywords Area
  - method
  - bayes_calibration
  - dream
  - emulator
  - pce
  - expansion_order
  - collocation_ratio
  - cross_validation
  - noise_only

Restrict the cross validation process to estimating only the best noise tolerance.
Specification

Alias: none
Argument(s): none
Default: false

Description

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.4.8.80 max_cv_order_candidates

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- cross_validation
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none
Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.
See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.4.8.81 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
7.2.68.4.8.82  response_scaling

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

Specification

Alias: none
Argument(s): none
Default: use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to $[0,1]$ based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`
- `multifidelity_function_train`

7.2.68.4.8.83  use_derivatives

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
Use derivative data to construct surrogate models

Specification

Alias: none
Argument(s): none
Default: use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.4.8.84 tensor_grid

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

Alias: none
Argument(s): none
Default: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
7.2.68.4.8.85 reuse_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.4.8.86 max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the
maximum iterations in the coefficient solver.

7.2.68.4.8.87 expansion_samples

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• pce
• expansion_order
• expansion_samples

Number of simulation samples used to estimate the expected value of a set of PCE coefficients

Specification

Alias: none
Argument(s): INTEGER

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td></td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

Description

In this case, the expansion order $p$ cannot be inferred from the numerical integration specification and it is necessary to provide an expansion_order to specify $p$ for a total-order expansion.

A corresponding sequence specification is documented at, e.g., expansion_samples_sequence and expansion_samples_sequence
See Also

These keywords may also be of interest:

- `expansion_samples_sequence` method-multilevel_polynomial_chaos-expansion_order_sequence-expansion_samples_sequence

7.2.68.4.8.88 reuse_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- expansion_samples
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

7.2.68.4.8.89 import_build_points_file

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- import_build_points_file

File containing points you wish to use to build a surrogate
**Specification**

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Selects custom-annotated tabular file format</td>
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<td></td>
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</tr>
<tr>
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<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.4.8.90 custom_annotated

- Keywords Area
- method
Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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</tr>
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<td>Enable evaluation ID column in custom-annotated tabular file</td>
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<td>interface_id</td>
<td>Dakota Keyword</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.
Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.4.8.91 header

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated
7.2.68.4.8.92  eval_id

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent custom_annotated

7.2.68.4.8.93  interface_id

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file
7.2 method

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.4.8.94 annotated

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- expansion_order
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
   tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.4.8.95  freeform

• **Keywords Area**

• **method**

• **bayes_calibration**

• **dream**

• **emulator**

• **pce**

• **expansion_order**

• **import_build_points_file**

• **freeform**

Selects freeform file format

Topics

This keyword is related to the topics:

• **file_formats**
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
0.9     1.1    0.0002    0.26     0.76
0.90009 1.1  0.0001996404857 0.2601620081 0.759955
0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...
```

Keywords Area

- method
- bayes_calibration
- dream
- emulator
- pce
import_build_points_file

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none
7.2 method

Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

Examples

bayes_calibration queso
  chain_samples = 2000 seed = 348
delayed_rejection
  emulator
    gaussian_process surfpack build_samples = 30
    posterior_adaptive max_iterations = 10
    proposal_covariance derivatives

7.2.68.4.8.98 orthogonal_least_interpolation

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>collocation_points</td>
<td></td>
<td>Number of collocation points used to estimate expansion coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td></td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>
### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

<table>
<thead>
<tr>
<th>7.2.68.4.8.99</th>
<th>collocation_points</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Keywords Area</strong></td>
<td></td>
</tr>
<tr>
<td><strong>method</strong></td>
<td></td>
</tr>
<tr>
<td><strong>bayes_calibration</strong></td>
<td></td>
</tr>
<tr>
<td><strong>dream</strong></td>
<td></td>
</tr>
<tr>
<td><strong>emulator</strong></td>
<td></td>
</tr>
<tr>
<td><strong>pce</strong></td>
<td></td>
</tr>
<tr>
<td><strong>orthogonal_least_interpolation</strong></td>
<td></td>
</tr>
<tr>
<td><strong>collocation_points</strong></td>
<td></td>
</tr>
</tbody>
</table>

Number of collocation points used to estimate expansion coefficients

### Specification

**Alias:** none

**Argument(s):** INTEGER

### Description

Specify the number of collocation points used to estimate expansion coefficients using regression approaches.

A corresponding sequence specification is documented at, e.g., `collocation_points_sequence` and `collocation_points_sequence`
7.2 method

See Also

These keywords may also be of interest:

- collocation_points_sequence
- method-multilevel_polynomial_chaos-expansion_order_sequence-collocation_ratio-collocation_points_sequence

7.2.68.4.8.100 tensor_grid

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias**: none

**Argument(s)**: INTEGERLIST

**Default**: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.4.8.101 reuse_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.
Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file
Argument(s): STRING
Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
</tr>
</tbody>
</table>
Optional | active_only | Import only active variables from tabular data file

Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.4.8.103 custom_annotated

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>custom_annotated</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>custom_annotated</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>custom_annotated</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:
<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.2.68.4.8.104  header

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no header

**Description**

See description of parent `custom_annotated`

7.2.68.4.8.105  eval_id

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
Specification

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

Description

See description of parent *custom.annotated*

7.2.68.4.8.106  interface_id

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **orthogonal_least_interpolation**
- **import_build_points_file**
- **custom.annotated**
- **interface_id**

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

Description

See description of parent *custom.annotated*

7.2.68.4.8.107  annotated

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **pce**
- **orthogonal_least_interpolation**
7.2 method

- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1   NO_ID   0.9  1.1  0.0002  0.26  0.76
2   NO_ID  0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3   NO_ID  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```
Selects freeform file format

Topics
This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```python
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>
```

7.2.68.4.8.109 active_only

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
Adapt emulator model to increase accuracy in high posterior probability regions

Specification

Alias: none
Argument(s): none

Description

Following an emulator-based MCMC process, this option refines the emulator by selecting points in regions of high posterior probability, performing truth evaluations at these points, updating the emulator, and reperforming the MCMC process. The adaptation is continued until the maximum number of iterations is exceeded or the convergence tolerance is met.

Examples

```plaintext
bayes_calibration queso
  chain_samples = 2000 seed = 348
  delayed_rejection
  emulator
    gaussian_process surfpack build_samples = 30
    posterior_adaptive max_iterations = 10
    proposal_covariance derivatives
```

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
7.2 method

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

7.2.68.4.8.112 wiener

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.4.8.113 normalized

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- normalized

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

Specification

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.4.8.114 export_expansion_file

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

Specification

**Alias:** none

**Argument(s):** STRING
Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consist only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.4.8.115 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- diagonal_covariance

Display only the diagonal terms of the covariance matrix

Specification

Alias: none
Argument(s): none
Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.4.8.116 full_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- pce
- full_covariance

Display the full covariance matrix

Specification

Alias: none
Argument(s): none
Description

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.68.4.8.117 ml_pce

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce

Multilevel Polynomial Chaos Expansion as an emulator model.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multilevel expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td>-----------------------</td>
<td>----------------------------------</td>
<td>-------</td>
<td>------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Optional</td>
<td>wiener</td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
<tr>
<td>Optional</td>
<td>normalized</td>
<td>normalized</td>
<td>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>export_expansion_file</td>
<td>export_expansion_file</td>
<td>Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 3)</td>
<td>diagonal_covariance</td>
<td>Display only the diagonal terms of the covariance matrix</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

### Description

Selects a multilevel polynomial chaos expansion (ML PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using ML PCE as a surrogate within the Bayesian framework.

### See Also

These keywords may also be of interest:

- `multilevel_polynomial_chaos`

- `allocation_control`
Sample allocation approach for multilevel expansions

**Specification**

Alias: none

Argument(s): none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Multilevel Sample Allocation Control (Group 1)</td>
<td>estimator_variance</td>
<td>Variance of mean estimator within multilevel polynomial chaos</td>
</tr>
<tr>
<td></td>
<td>rip_sampling</td>
<td></td>
<td>Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

**Description**

Multilevel expansions, including regression-based polynomial chaos expansion (PCE) and function train (FT) expansions, require a sample allocation strategy. Three options are currently available:

- allocation based on assuming a convergence rate for the estimator variance (for regression PCE)
- restricted isometry property (RIP) sampling (for regression PCE via compressed sensing)
- rank sampling (for FT)

**Default Behavior**

Current defaults, when allocation_control is not specified, are estimator variance for PCE and rank sampling for FT.

7.2.68.4.8.119 estimator_variance
• emulator
• ml_pce
• allocation_control
• estimator_variance

Variance of mean estimator within multilevel polynomial chaos

Specification

Alias: none
Argument(s): none
Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>estimator_rate</td>
<td>Rate of convergence of mean estimator within multilevel polynomial chaos</td>
</tr>
</tbody>
</table>

Description

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
\text{Var}[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}
\]

for free parameters \(\gamma\) and \(\kappa\), with default values that may be overridden as part of this specification block.

This approach is supported for regression-based PCE approaches (over-determined least squares, under-determined compressed sensing, or orthogonal least interpolation).

In practice, it can be challenging to estimate a smooth convergence rate for estimator variance in the presence of abrupt transitions in the quality of sparse recoveries. As a result, sample allocation by greedy refinement is generally preferred.

Examples

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the convergence of the estimator variance, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.
Rate of convergence of mean estimator within multilevel polynomial chaos

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 2

**Description**

Multilevel Monte Carlo performs optimal resource allocation based on a known estimator variance for the mean statistic:

\[
Var[\hat{Q}] = \frac{\sigma_Q^2}{N}
\]

Replacing the simple ensemble average estimator in Monte Carlo with a polynomial chaos estimator results in a different and unknown relationship between the estimator variance and the number of samples. In one approach to multilevel PCE, we can employ a parameterized estimator variance:

\[
Var[\hat{Q}] = \frac{\sigma_Q^2}{\gamma N^\kappa}
\]

for free parameters \(\gamma\) and \(\kappa\).

The default values are \(\gamma = 1\) and \(\kappa = 2\) (adopts a more aggressive sample profile by assuming a faster convergence rate than Monte Carlo). This advanced specification option allows to user to specify \(\kappa\), overriding the default.
Sample allocation based on restricted isometry property (RIP) within multilevel polynomial chaos

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Multilevel polynomial chaos with compressed sensing may allocate the number of samples per level based on the restricted isometry property (RIP), applied to recovery at level \( l \):

\[ N_l \geq s_l \log^3(s_l) L_l \log(C_l) \]

for sparsity \( s_l \), cardinality \( C_l \), and mutual coherence \( L_l \). The adaptive algorithm starts from a pilot sample, shapes the profile based on observed sparsity, and iterates until convergence. In practice, RIP sampling levels are quite conservative, and a collocation ratio constraint (\( N_l \leq rC_l \), where \( r \) defaults to 2) must be enforced on the profile.

The algorithm relies on observed sparsity, it is appropriate for use with regularized solvers for compressed sensing. It employs orthogonal matching pursuit (OMP) by default and automatically activates cross-validation in order to choose the best noise parameter value for the recovery.

This capability is **experimental**. Sample allocation by greedy refinement is generally preferred.

**Examples**

This example starts with sparse recovery for a second-order candidate expansion at each level. As the number of samples is adapted for each level, as dictated by the number of sparse coefficient sets recovered for each level, the candidate expansion order is incremented as needed in order to synchronize with the specified collocation ratio.

```plaintext
method, model_pointer = 'HIERARCH'
   multilevel_polynomial_chaos
      orthogonal_matching_pursuit
         expansion_order_sequence = 2
         pilot_samples = 10
         collocation_ratio = .9
         allocation_control rip_sampling
         seed = 1237
         convergence_tolerance = .01
```
In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the **distinct** approach may tend to be more expensive in exchange for greater robustness.
7.2 method

Examples

```
method,
  multilevel_polynomial_chaos
evaluation_order_sequence = 2
collocation_ratio = .9
orthogonal_matching_pursuit
discrepancy_emulation recursive
```

7.2.68.4.8.123  distinct

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- discrepancy_emulation
- distinct

Distinct formulation for emulation of model discrepancies.

Specification

Alias: paired

Argument(s): none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.4.8.124  recursive

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- discrepancy_emulation
- recursive

Recursive formulation for emulation of model discrepancies.
Specification

**Alias:** none

**Argument(s):** none

**Description**

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

**Examples**

7.2.68.4.8.125  expansion_order_sequence

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**

Sequence of expansion orders used in a multi-stage expansion

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>----------------------</td>
<td>---------</td>
<td></td>
</tr>
<tr>
<td><strong>basis_type</strong></td>
<td><strong>collocation_ratio</strong></td>
<td><strong>expansion_samples_sequence</strong></td>
<td></td>
</tr>
<tr>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the <code>expansion_samples_sequence</code> applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., <code>expansion_samples</code></td>
<td></td>
</tr>
</tbody>
</table>
**Import-Build-Points File**

**Optional**

| import_build_points_file | File containing points you wish to use to build a surrogate |

**Description**

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`.

**See Also**

These keywords may also be of interest:

- `expansion_order`

7.2.68.4.8.126  **Dimension Preference**

| Keywords Area |
| method |
| bayes_calibration |
| dream |
| emulator |
| ml_pce |
| expansion_order_sequence |
| dimension_preference |

A set of weights specifying the relative importance of each uncertain variable (dimension).

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.6.8.4.127 basis_type

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- basis_type

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<th>Required/Optional</th>
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<td>Required (Choose One)</td>
<td>PCE Basis Type (Group 1)</td>
<td>tensor_product</td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>total_order</td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapted</td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

Description

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order \( p \) keep all terms with \( d \)-dimensional multi index \( i = (i_1, \ldots, i_d) \) that satisfies

\[
\max (i_1, \ldots, i_d) \leq p
\]

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none
Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order \( p \) keep all terms with a \( d \)-dimensional multi index \( \mathbf{i} = (i_1, \ldots, i_d) \) that satisfies

\[
\sum_{k=1}^{d} i_k \leq p
\]

7.2.68.4.8.130 adapted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>advancements</td>
<td></td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td>soft_convergence_limit</td>
<td></td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2.68.4.8.131 advancements

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted
- advancements

The maximum number of steps used to expand a basis step.

Specification

**Alias:** none

**Argument(s):** INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non-zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.4.8.132 soft_convergence_limit

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- basis_type
- adapted
- soft_convergence_limit

The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.
7.2 method

Specification

Alias: none
Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify an initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.4.8.133  collocation_ratio

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none
Argument(s): REAL
Child Keywords:

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<tr>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Regression Algorithm (Group 1)</td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
<tr>
<td><strong>orthogonal_matching_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
<td></td>
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</tr>
<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td><strong>cross_validation</strong></td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td><strong>ratio_order</strong></td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
<td></td>
</tr>
</tbody>
</table>
### 7.2 method

<table>
<thead>
<tr>
<th>Optional</th>
<th>response_scaling</th>
<th>Perform bounds-scaling on response values prior to surrogate emulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_solver_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
</tr>
</tbody>
</table>

### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \(cN^o\) where \(c\) is the collocation_ratio and \(o\) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \(cN^{o+1}\)).

### collocation_points_sequence

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion
Specification

Alias: pilot_samples

Argument(s): INTEGERLIST

Description

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

See Also

These keywords may also be of interest:

- `collocation_points`

7.2.68.4.135 least_squares

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

Alias: none

Argument(s): none

Default: svd

Child Keywords:
### Description

Compute the coefficients of a polynomial expansion using least squares. Specifically, SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

### Specification

- **Alias**: none
- **Argument(s)**: none

### Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

---

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias:** omp  
**Argument(s):** none  
**Child Keywords:**
Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP). Orthogonal matching pursuit (OMP) is a greedy algorithm that is useful when solving underdetermined linear systems.

7.2.68.4.8.139 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.140 basis_pursuit

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.4.8.141 basis_pursuit_denoising

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• ml_pce
• expansion_order_sequence
• collocation_ratio
• basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>
Description

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

7.2.68.4.8.142 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

**Alias:** none

**Argument(s):** REALIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.143 least_angle_regression

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
Specification

Alias: lars

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

Description

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.4.8.144 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

Specification

Alias: none

Argument(s): REALLIST

Default: 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

Description

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso  
**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.147  l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_only</td>
<td></td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td>max_cv_order_candidates</td>
<td></td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** false  

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.4.8.150 max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** ushort max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.4.8.151 ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none

Argument(s): REAL

Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^{o}$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
7.2.68.4.8.152 response_scaling

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- multifidelity_polynomial_chaos
- multifidelity_function_train

7.2.68.4.8.153 use_derivatives

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
Use derivative data to construct surrogate models

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
7.2.68.4.8.155 reuse_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.4.8.156 max_solver_iterations

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- collocation_ratio
- max_solver_iterations

Maximum iterations in determining polynomial coefficients
### 7.2 method

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

**7.2.68.4.8.157 expansion_samples_sequence**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **ml_pce**
- **expansion_order_sequence**
- **expansion_samples_sequence**

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the `expansion_samples_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_samples`

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>

**See Also**

These keywords may also be of interest:

- `expansion_samples`
7.2.68.4.8.158  reuse_points

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- expansion_samples_sequence
- reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples

Argument(s): none

Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

7.2.68.4.8.159  import_build_points_file

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- import_build_points_file

File containing points you wish to use to build a surrogate
Specification

**Alias:** import_points_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

#### Default Behavior

By default, methods do not import points from a file.

#### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

#### Examples

```dakota
method
  polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### Examples

- `Keywords Area`
- `method`
- `bayes_calibration`
- `dream`
- `emulator`
- `ml_pce`
- `expansion_order_sequence`
• import_build_points_file
• custom_annotated

Selects custom-annotated tabular file format

Topics
This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
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<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id

Resulting tabular file:

%eval_id x1       x2       obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1     0.0002  0.26       0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.4.8.161 header

• Keywords Area
  • method
  • bayes_calibration
  • dream
  • emulator
  • ml_pce
  • expansion_order_sequence
  • import_build_points_file
  • custom_annotated
  • header

Enable header row in custom-annotated tabular file

Specification

 Alias: none
 Argument(s): none
 Default: no header

Description

See description of parent custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent custom.annotated
**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`  
7.2.68.4.8.164  annotated  

- **Keywords Area**  
- method  
- bayes_calibration  
- dream  
- emulator  
- ml_pce  
- expansion_order_sequence  
- import_build_points_file  
- annotated

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```environment
  tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.4.165 freeform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- expansion_order_sequence
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
 0.9  1.1  0.0002  0.26  0.76
 0.90009  1.1  0.0001996404857  0.2601620081  0.759955
 0.89991  1.1  0.0002003604863  0.2598380081  0.760045
```

7.2.68.4.8.166  active_only

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
• expansion_order_sequence
• import_build_points_file
• active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword active_only indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.4.8.167 orthogonal_least_interpolation

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• ml_pce
• orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

Alias: least_interpolation oli
Argument(s): none
Child Keywords:
## Keywords Area

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
</tbody>
</table>

### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

#### 7.2.68.4.8.168  collocation_points_sequence

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- orthogonal_least_interpolation
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion.

### Specification

**Alias**: none

**Argument(s)**: INTEGERLIST
Description

Each level entry of the \texttt{collocation_points_sequence} applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., \texttt{collocation_points}

See Also

These keywords may also be of interest:

- \texttt{collocation_points}

7.2.68.4.8.169 tensor_grid

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{bayes_calibration}
- \texttt{emulator}
- \texttt{ml_pce}
- \texttt{orthogonal_least_interpolation}
- \texttt{tensor_grid}

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: INTEGERLIST

\textbf{Default}: regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.2.68.4.8.170 reuse_points

- \texttt{Keywords Area}
- \texttt{method}
- \texttt{bayes_calibration}
- \texttt{dream}
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

**Specification**

**Alias:** reuse_samples  
**Argument(s):** none  
**Default:** no sample reuse in coefficient estimation

**Description**

The `reuse_points` option controls the reuse behavior of points for various types of polynomial chaos expansions, including: `collocation_points`, `collocation_ratio`, `expansion_samples`, or `orthogonal_least_interpolation`. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify `reuse_points` so that any points that have been previously generated (for example, from the `import_points_file`) can be reused.

**7.2.68.4.8.171 import_build_points_file**

- **Keywords Area**  
- **method**  
- **bayes_calibration**  
- **dream**  
- **emulator**  
- **ml_pce**  
- **orthogonal_least_interpolation**  
- **import_build_points_file**

File containing points you wish to use to build a surrogate

**Specification**

**Alias:** import_points_file  
**Argument(s):** STRING  
**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>
### Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

### Default Behavior

By default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

### Examples

```dakota
method polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

### Custom annotated

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
**custom_annotated**

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- `file_formats`

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether `header` row, `eval_id` column, and `interface_id` column appear in the tabular file, thus bridging `freeform` and (fully) annotated.

**Default Behavior**

The `annotated` format is the default for tabular export/import. To control which header row and columns are in the input/output, specify `custom_annotated`, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1  x2  obj_fn  nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.4.8.173 header

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.68.4.8.174 eval_id

- Keywords Area
- method
- bayes_calibration
- dream
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated_header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```
%eval_id interface  x1   x2    obj_fn nln_ineq_con_1 nln_ineq_con_2
1     NO_ID        0.9   1.1   0.0002   0.26    0.76
2     NO_ID        0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3     NO_ID        0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...```

```
7.2.68.4.8.177 freeform
    ```
    • Keywords Area
    • method
    • bayes_calibration
    • dream
    • emulator
    • ml_pce
    • orthogonal_least_interpolation
    • import_build_points_file
    • freeform
    ```
```

Selects freeform file format

Topics

This keyword is related to the topics:

```
• file_formats
```

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tenabular_data
   tabular_data_file = 'dakota_summary.dat'
   freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.4.8.178  active_only

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- orthogonal_least_interpolation
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Description**

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.4.8.179 `askey`

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **ml_pce**
- **askey**

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

**Description**

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

**See Also**

These keywords may also be of interest:

- **polynomial_chaos**
- **wiener**
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias**: none  
**Argument(s)**: none  
**Default**: extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- `polynomial_chaos`
- `askey`

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
Specification

Alias: none
Argument(s): none
Default: PCE coefficients correspond to unnormalized basis polynomials

Description

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.

7.2.68.4.8.182 export_expansion_file

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- export_expansion_file

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.

Specification

Alias: none
Argument(s): STRING

Description

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.4.8.183 diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- ml_pce
- diagonal_covariance

Display only the diagonal terms of the covariance matrix.
7.2 method

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** diagonal_covariance for response vector > 10; else full_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. *diagonal_covariance* is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

```
7.2.68.4.8.184  full_covariance
```

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **ml_pce**
- **full_covariance**

Display the full covariance matrix

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. *full_covariance* is used to force Dakota to output the full covariance matrix.

```
7.2.68.4.8.185  mf_pce
```

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**

Multifidelity Polynomial Chaos Expansion as an emulator model.
Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**: none
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_refinement_order_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>allocation_control</td>
<td>Sample allocation approach for multifidelity expansions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrepancy_emulation</td>
<td>Formulation for emulation of model discrepancies.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>quadrature_order_sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sparse_grid_level_sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>expansion_order_sequence</td>
<td>Sequence of expansion orders used in a multi-stage expansion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>orthogonal_least_interpolation</td>
<td>Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 2)</td>
<td>askkey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>
The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials. Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.

<table>
<thead>
<tr>
<th>Optional</th>
<th>normalized</th>
<th>The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials. Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>export_expansion_file</td>
<td>Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Covariance Type (Group 3)</td>
<td>diagonal_covariance</td>
</tr>
<tr>
<td></td>
<td>full_covariance</td>
<td>Display the full covariance matrix</td>
</tr>
</tbody>
</table>

**Description**

Selects a multifidelity polynomial chaos expansion (MF PCE) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF PCE as a surrogate within the Bayesian framework.

**See Also**

These keywords may also be of interest:

- multifidelity_polynomial_chaos

**Automatic polynomial order refinement**

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: no refinement

**Child Keywords**: none
### 7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group p-refinement Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>uniform</td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
</tbody>
</table>

**Description**

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol’ indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.4.187 uniform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

**7.2.68.4.8.188 dimension_adaptive**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **p_refinement**
- **dimension_adaptive**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.  

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

#### 7.2.68.4.8.189 sobol

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.
Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```
method,
   polynomial_chaos
   sparse_grid_level = 3
   dimension_adaptive p_refinement sobol
   max_iterations = 20
   convergence_tol = 1.e-4
```

7.2.68.4.8.190 decay

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• p_refinement
• dimension_adaptive
• decay

Estimate spectral coefficient decay rates to guide dimension-adaptive refinement.

Specification

Alias: none
Argument(s): none

Description

Estimate spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation. These decay rates are used to guide dimension-adaptive refinement, where slower decay rates result in higher dimension preference.
Examples

```
method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive p_refinement decay
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.4.191 generalized

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- p_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

**Alias:** none

**Argument(s):** none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```
method, polynomial_chaos
  sparse_grid_level = 3
dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```

7.2.68.4.192 max_refinement_iterations

- Keywords Area
- method
- bayes_calibration
Maximum number of expansion refinement iterations

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.

**Examples**

```
method, polynomial_chaos
  dimension_adaptive generalized
p_refinement
  max_refinement_iterations = 20
  convergence_tol = 1.e-4
  sparse_grid_level = 1
```

**Sample allocation approach for multifidelity expansions**

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

- allocation_control
# Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ an integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

## Default Behavior

The default, when `allocation_control` is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).

### 7.2.68.4.8.194 greedy

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **allocation_control**
- **greedy**

Sample allocation based on greedy refinement within multifidelity polynomial chaos

## Specification

**Alias:** none

**Argument(s):** none

## Description

Multifidelity polynomial chaos supports greedy refinement strategies, spanning regression and projection approaches for computing expansion coefficients. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within a unified competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any \( z/p/\beta/\beta^* \) level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.
Examples

The following example of greedy multifidelity regression starts from a zeroth-order reference expansion (a constant) for each level, and generates candidate refinements for each level that are competed in an integrated greedy competition. The number of new samples for the incremented candidate expansion order is determined from the collocation ratio. In this case, the number of candidates for each level is limited to one uniform refinement of the current expansion order.

```
method,
    model_pointer = 'HIERARCH'
    multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement uniform
    expansion_order_sequence = 0
    collocation_ratio = .9  seed = 160415
    orthogonal_matching_pursuit
    convergence_tolerance 1.e-2
```

The next example employs generalized sparse grids within a greedy multifidelity competition. Each modeling level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all model levels is competed within an integrated greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each model level and the number of candidates grows rapidly with random dimension and grid level.

```
method,
    model_pointer = 'HIERARCH'
    multifidelity_polynomial_chaos
    allocation_control greedy
    p_refinement dimension_adaptive generalized
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-8
```

7.2.68.4.8.195 discrepancy_emulation

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• discrepancy_emulation

Formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none
Default: distinct
Child Keywords:
Description

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. distinct emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.

2. recursive emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.

The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

Examples

```
method,
    multilevel_polynomial_choas
    expansion_order_sequence = 2
    collocation_ratio = .9
    orthogonal_matching_pursuit
    discrepancy_emulation recursive
```

```
7.2.68.4.8.196  distinct
```

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- discrepancy_emulation
- distinct

Distinct formulation for emulation of model discrepancies.
Specification

Alias: paired
Argument(s): none

Description

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

Examples

7.2.68.4.8.197 recursive

Recursive formulation for emulation of model discrepancies.

Specification

Alias: none
Argument(s): none

Description

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

Examples
7.2.68.4.8.198 quadrature_order_sequence

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- quadrature_order_sequence

Sequence of quadrature orders used in a multi-stage expansion

**Specification**

**Alias**: none

**Argument(s)**: INTEGERLIST

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td></td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional/Choose One</td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

**Description**

Each level entry of the quadrature_order_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., quadrature_order

**See Also**

These keywords may also be of interest:

- quadrature_order
7.2.68.4.8.199  dimension_preference

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- quadrature_order_sequence
- dimension_preference

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leada to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.200  nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available.
Specification

Alias: none
Argument(s): none
Default: quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.201 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules

Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.4.8.202 sparse_grid_level_sequence

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion
Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Growth (Group 1)</td>
<td>restricted</td>
<td>Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>unrestricted</td>
<td>Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Quadrature Rule Nesting (Group 2)</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td></td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

**Description**

Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`

**See Also**

These keywords may also be of interest:

- `sparse_grid_level`

7.2.68.4.8.203  `dimension_preference`

- Keywords Area
- method
A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.204  restricted

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.205 unrestricted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

**Alias:** none

**Argument(s):** none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
Enforce use of nested quadrature rules if available

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

Enforce use of non-nested quadrature rules

**Specification**

**Alias:** none

**Argument(s):** none
Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.4.8.208  expansion_order_sequence

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• expansion_order_sequence

Sequence of expansion orders used in a multi-stage expansion

Specification

Alias: none

Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>basis_type</td>
<td>Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Group 1</td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
<tr>
<td><strong>expansion_samples_sequence</strong></td>
<td>Sequence of expansion samples used in a multi-stage polynomial chaos expansion. Each level entry of the <code>expansion_samples_sequence</code> applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos. If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach. A corresponding scalar specification is documented at, e.g., <code>expansion_samples</code></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### dimension_preference

**Description**

Each level entry of the `expansion_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion order sequences include multilevel or multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `expansion_order`

**See Also**

These keywords may also be of interest:

- `expansion_order`

**Specification**

- **Alias:** none
- **Argument(s):** REALLIST
- **Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- `sobol`
- `decay`

### 7.2.68.4.8.210 basis_type

- **Keywords Area**
- `method`
- `bayes_calibration`
- `dream`
- `emulator`
- `mf_pce`
- `expansion_order_sequence`
- `basis_type`

Specify the type of basis truncation to be used for a Polynomial Chaos Expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Description of Group PCE Basis Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td><code>tensor_product</code></td>
<td>Use a tensor-product index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>total_order</code></td>
<td>Use a total-order index set to construct a polynomial chaos expansion.</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>adapted</code></td>
<td>Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.</td>
</tr>
</tbody>
</table>

**Description**

Specify the type of basis truncation that determines the terms that are included within a Polynomial Chaos Expansion. Options include tensor-product, total-order, and adapted.
Use a tensor-product index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Use a tensor-product index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with $d$-dimensional multi index $i = (i_1, \ldots, i_d)$ that satisfies

$$\max(i_1, \ldots, i_d) \leq p$$

Use a total-order index set to construct a polynomial chaos expansion.

**Specification**

**Alias**: none

**Argument(s)**: none
Description

Use the traditional total-order index set to construct a polynomial chaos expansion. That is for a given order $p$ keep all terms with a $d$-dimensional multi index $\mathbf{i} = (i_1, \ldots, i_d)$ that satisfies

$$\sum_{k=1}^{d} i_k \leq p$$

7.2.68.4.8.213 adapted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- basis_type
- adapted

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion.

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>advancements</td>
<td></td>
<td>The maximum number of steps used to expand a basis step.</td>
</tr>
<tr>
<td>Optional</td>
<td>soft_convergence_limit</td>
<td></td>
<td>The maximum number of times no improvement in cross validation error is allowed before the algorithm is terminated.</td>
</tr>
</tbody>
</table>

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
The maximum number of steps used to expand a basis step.

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.
7.2 method

Specification

Alias: none
Argument(s): INTEGER

Description

Use adaptive basis selection to choose the basis terms in a polynomial chaos expansion. Basis selection uses compressed sensing to identify a initial set of non zero PCE coefficients. Then these non-zero terms are expanded a set number of times (we suggest 3) and compressed sensing is then applied to these three new index sets. Cross validation is used to choose the best candidate basis. The best basis is then restricted again to the non-zero terms and expanded until no improvement can be gained by adding additional terms.

7.2.68.4.8.216 collocation_ratio

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none
Argument(s): REAL

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>collocation_points_sequence</td>
<td>Sequence of collocation point counts used in a multi-stage expansion</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>least_squares</td>
<td>Compute the coefficients of a polynomial expansion using least squares</td>
</tr>
</tbody>
</table>

| Regression Algorithm (Group 1) |

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<table>
<thead>
<tr>
<th><strong>orthogonal_matching_pursuit</strong></th>
<th>Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>basis_pursuit</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.</td>
</tr>
<tr>
<td><strong>basis_pursuit_denoising</strong></td>
<td>Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.</td>
</tr>
<tr>
<td><strong>least_angle_regression</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.</td>
</tr>
<tr>
<td><strong>least_absolute_shrinkage</strong></td>
<td>Compute the coefficients of a polynomial expansion by using the LASSO problem.</td>
</tr>
<tr>
<td><strong>cross_validation</strong></td>
<td>Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.</td>
</tr>
<tr>
<td><strong>ratio_order</strong></td>
<td>Specify a non-linear relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.</td>
</tr>
</tbody>
</table>
### Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the collocation_ratio allows for specification of a constant factor applied to \( N \) (e.g., collocation_ratio = 2. produces samples = 2\( N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \( cN^o \) where \( c \) is the collocation_ratio and \( o \) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( cN^o \frac{n}{p+1} \)).

#### 7.2.68.4.8.217 collocation_points_sequence

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **expansion_order_sequence**
- **collocation_ratio**
- **collocation_points_sequence**

Sequence of collocation point counts used in a multi-stage expansion
Specification

Alias: pilot_samples
Argument(s): INTEGERLIST

Description

Each level entry of the collocation_points_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., collocation_points

See Also

These keywords may also be of interest:

- collocation_points

7.2.68.4.218 least_squares

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_squares

Compute the coefficients of a polynomial expansion using least squares

Specification

Alias: none
Argument(s): none
Default: svd

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Compute the coefficients of a polynomial expansion using least squares. Specifically, SVD-based least-squares will be used for solving over-determined systems. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares.

### Specification

**Alias:** none

**Argument(s):** none

### Description

Calculate the coefficients of a polynomial chaos expansion using the singular value decomposition. When the number of model runs exceeds the number of terms in the PCE, the solution returned will be the least-squares solution, otherwise the solution will be the minimum norm solution computed using the pseudo-inverse.
Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Calculate the coefficients of a polynomial chaos expansion using equality constrained least squares.

**Specification**

**Alias**: omp

**Argument(s)**: none

**Child Keywords:**

7.2.68.4.8.220  equality_constrained

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• expansion_order_sequence
• collocation_ratio
• least_squares
• equality_constrained

Compute the coefficients of a polynomial expansion using orthogonal matching pursuit (OMP)

**Specification**

**Alias**: omp

**Argument(s)**: none

**Child Keywords:**
### 7.2.68.4.8.222 noise_tolerance

**Keywords Area**

- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- orthogonal_matching_pursuit
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

#### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

### 7.2.68.4.8.223 basis_pursuit

**Keywords Area**

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
• mf_pce
• expansion_order_sequence
• collocation_ratio
• basis_pursuit

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

**Specification**

**Alias:** bp

**Argument(s):** none

**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit $\ell_1$-minimization problem using linear programming.

7.2.68.4.8.224 basis_pursuit_denoising

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• expansion_order_sequence
• collocation_ratio
• basis_pursuit_denoising

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**Specification**

**Alias:** bpdn

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td>noise_tolerance</td>
<td></td>
</tr>
</tbody>
</table>

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
**Description**

Compute the coefficients of a polynomial expansion by solving the Basis Pursuit Denoising $\ell_1$-minimization problem using second order cone optimization.

**7.2.68.4.8.225 noise_tolerance**

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- basis_pursuit_denoising
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**7.2.68.4.8.226 least_angle_regression**

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.
**Specification**

**Alias:** lars  
**Argument(s):** none  

**Child Keywords:**

<table>
<thead>
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<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_tolerance</td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors.</td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the greedy least angle regression (LAR) method.

7.2.68.4.8.227 noise_tolerance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- least_angle_regression
- noise_tolerance

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  
**Argument(s):** REALLIST  
**Default:** 1e-3 for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.
Compute the coefficients of a polynomial expansion by using the LASSO problem.

**Specification**

**Alias:** lasso  
**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>noise_tolerance</td>
<td></td>
<td>The noise tolerance used when performing cross validation in the presence of noise or truncation errors. The $l_2$ penalty used when performing compressed sensing with elastic net.</td>
</tr>
<tr>
<td>Optional</td>
<td>l2_penalty</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Compute the coefficients of a polynomial expansion by using the LASSO problem.
The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

**Specification**

**Alias:** none  

**Argument(s):** REALLIST  

**Default:** $1e^{-3}$ for BPDN, 0. otherwise (algorithms run until termination)

**Description**

The noise tolerance used when performing cross validation in the presence of noise or truncation errors.

7.2.68.4.8.230  l2_penalty

The $l_2$ penalty used when performing compressed sensing with elastic net.

**Specification**

**Alias:** none  

**Argument(s):** REAL  

**Default:** 0. (reverts to standard LASSO formulation)

**Description**

The $l_2$ penalty used when performing compressed sensing with elastic net.
Use cross validation to choose the 'best' polynomial order of a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>noise_only</td>
<td>Restrict the cross validation process to estimating only the best noise tolerance.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order</td>
</tr>
</tbody>
</table>

**Description**

Use cross validation to choose the 'best' polynomial degree of a polynomial chaos expansion. 10 fold cross validation is used to estimate the cross validation error of a total-order polynomial expansion for orders 1 through to order. The order chosen is the one that produces the lowest cross validation error. If there are not enough points to perform 10 fold cross validation then one-at-a-time cross validation will be performed.
Restrict the cross validation process to estimating only the best noise tolerance.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

By default, cross validation estimates both the best noise tolerance and the best candidate basis order. For reasons of reducing computational cost by reducing the number of candidate solves, the user may wish to restrict this process to estimating only the best noise tolerance.

Generally speaking, computing the best noise tolerance through cross validation mitigates issues with over-fitting the data. Computing the best candidate basis order can also mitigate overfitting, while also controlling levels of mutual coherence resulting from high-order Vandermonde-like matrix systems.

7.2.68.4.8.233 max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** ushort max
Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.2.68.4.8.234 ratio_order

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
- collocation_ratio
- ratio_order

Specify a non-linear the relationship between the expansion order of a polynomial chaos expansion and the number of samples that will be used to compute the PCE coefficients.

Specification

Alias: none
Argument(s): REAL
Default: 1.

Description

When using regression type methods (specified with either collocation_points or collocation_ratio), a total-order expansion can be specified using expansion_order. To avoid requiring the user to calculate N from n and p), the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2. produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes $cN^o$ where $c$ is the collocation_ratio and $o$ is the ratio_order.
Perform bounds-scaling on response values prior to surrogate emulation

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** use original data

**Description**

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating `response_scaling`, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

**See Also**

These keywords may also be of interest:

- `multifidelity_polynomial_chaos`  
- `multifidelity_function_train`
Use derivative data to construct surrogate models

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

---

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** regression with LHS sample set (point collocation)

**Description**

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
reuse_points

Keywords Area
method
bayes_calibration
dream
emulator
mf_pce
expansion_order_sequence
collocation_ratio
reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

max_solver_iterations

Keywords Area
method
bayes_calibration
dream
emulator
mf_pce
expansion_order_sequence
collocation_ratio
max_solver_iterations

Maximum iterations in determining polynomial coefficients
Specification

Alias: none
Argument(s): INTEGER
Default: 100

Description

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.2.68.4.8.240 expansion_samples_sequence

Sequence of expansion samples used in a multi-stage polynomial chaos expansion

Each level entry of the expansion_samples_sequence applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support expansion samples sequences include multilevel and multifidelity polynomial chaos.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., expansion_samples

Specification

Alias: none
Argument(s): INTEGERLIST

Child Keywords:

```
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>reuse_points</td>
<td>This describes the behavior of reuse of points in constructing polynomial chaos expansion models.</td>
</tr>
</tbody>
</table>
```

See Also

These keywords may also be of interest:

- expansion_samples
This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

Alias: reuse_samples
Argument(s): none
Default: no sample reuse in coefficient estimation

Description

The reuse_points option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify reuse_points so that any points that have been previously generated (for example, from the import_points file) can be reused.

File containing points you wish to use to build a surrogate
Specification

**Alias**: import_points_file

**Argument(s)**: STRING

**Default**: no point import from a file

**Child Keywords**: 
Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```plaintext
method
  polynomial Chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

7.2.68.4.8.243 custom_annotated

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- expansion_order_sequence
7.2 method

- `import_build_points_file`
- `custom_annotated`

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- `file_formats`

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The `annotated` format is the default for tabular export/import. To control which header row and columns are in the input/output, specify `custom_annotated`, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.10002123457  0.26000234567  0.759955
3  0.89999  1.10002123457  0.2598380081  0.760045
...```

7.2.68.4.8.244 header

• Keywords Area

• method

• bayes_calibration

• dream

• emulator

• mf_pce

• expansion_order_sequence

• import_build_points_file

• custom_annotated

• header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Default: no header

Description

See description of parent custom_annotated
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated

7.2.68.4.8.247 annotated

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• expansion_order_sequence
• import_build_points_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips
To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

Selects freeform file format

Keywords Area

method

bayes_calibration

dream

emulator

mf_pce

expansion_order_sequence

import_build_points_file

freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none
Argument(s): none
Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = "dakota_summary.dat"
freeform

Resulting tabular file:

| 0.9  | 1.1  | 0.0002 | 0.26  | 0.76  |
| 0.90009 | 1.1 | 0.0001996404857 | 0.2601620081 | 0.759955 |
| 0.89991 | 1.1 | 0.0002003604863 | 0.2598380081 | 0.760045 |

7.2.68.4.8.249 active_only
```

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
7.2 method

- expansion_order_sequence
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Description**

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.4.8.250 orthogonal_least_interpolation

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- orthogonal_least_interpolation

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation.

Specification

**Alias:** least_interpolation oli

**Argument(s):** none

**Child Keywords:**
### Description

Build a polynomial chaos expansion from simulation samples using orthogonal least interpolation. Unlike the other regression methods `expansion_order` cannot be set. OLI will produce the lowest degree polynomial that interpolates the data.

#### 7.2.68.4.8.251 collocation_points_sequence

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- orthogonal_least_interpolation
- collocation_points_sequence

Sequence of collocation point counts used in a multi-stage expansion

### Specification

**Alias:** none

**Argument(s):** INTEGERLIST
Description

Each level entry of the `collocation_points_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support collocation point sequences include multilevel and multifidelity polynomial chaos and multilevel and multifidelity function train expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `collocation_points`

See Also

These keywords may also be of interest:

- `collocation_points`

7.2.68.4.8.252 tensor_grid

- `Keywords Area`
- `method`
- `bayes_calibration`
- `emulator`
- `mf_pce`
- `orthogonal_least_interpolation`
- `tensor_grid`

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** regression with LHS sample set (point collocation)

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.
• emulator
• mf_pce
• orthogonal_least_interpolation
• reuse_points

This describes the behavior of reuse of points in constructing polynomial chaos expansion models.

Specification

**Alias**: reuse_samples

**Argument(s)**: none

**Default**: no sample reuse in coefficient estimation

**Description**

The **reuse_points** option controls the reuse behavior of points for various types of polynomial chaos expansions, including: collocation_points, collocation_ratio, expansion_samples, or orthogonal_least_interpolation. If any of these approaches are specified to create a set of points for the polynomial chaos expansion, one can specify **reuse_points** so that any points that have been previously generated (for example, from the **import_points_file**) can be reused.

7.2.68.4.8.254  import_build_points_file

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_pce
• orthogonal_least_interpolation
• import_build_points_file

File containing points you wish to use to build a surrogate

Specification

**Alias**: import_points_file

**Argument(s)**: STRING

**Default**: no point import from a file

**Child Keywords**:
The import_build_points_file allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**Examples**

```plaintext
method
polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'
```

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2    obj_fn  nln_ineq_con_1  nln_ineq_con_2
1       0.9  1.1    0.0002  0.26         0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.4.8.256  header

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

**Default:** no header

Description

See description of parent custom_annotated

7.2.68.4.8.257  eval_id

- Keywords Area
- method
- bayes_calibration
- dream
Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no eval_id column

**Description**

See description of parent `custom_annotated`

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no interface_id column

**Description**

See description of parent `custom_annotated`
Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```python
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1  NO_ID   0.9  1.1  0.0002  0.26   0.76
2  NO_ID   0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3  NO_ID   0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.4.8.260  freeform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- orthogonal_least_interpolation
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009 1.1 0.0001996404857 0.2601620081 0.759955
  0.89991 1.1 0.0002003604863 0.2598380081 0.760045
  ...
```

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats
Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword *active_only* indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.68.4.8.262 askey

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_pce
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener
Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

**Description**

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).

**See Also**

These keywords may also be of interest:

- polynomial_chaos
- askey

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials.
Specification

**Alias:** none

**Argument(s):** none

**Default:** PCE coefficients correspond to unnormalized basis polynomials

**Description**

The normalized specification requests output of PCE coefficients that correspond to normalized orthogonal basis polynomials

7.2.68.4.8.265  export_expansion_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **export_expansion_file**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file

**Specification**

**Alias:** none

**Argument(s):** STRING

**Description**

Export the coefficients and multi-index of a Polynomial Chaos Expansion (PCE) to a file. The multi-index written will be sparse. Specifically the expansion will consists only of the indices corresponding to the non-zero coefficients of the PCE.

7.2.68.4.8.266  diagonal_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **diagonal_covariance**

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal_covariance for response vector $> 10$; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. `diagonal_covariance` is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.4.8.267 full_covariance

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_pce**
- **full_covariance**

Display the full covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. `full_covariance` is used to force Dakota to output the full covariance matrix.

7.2.68.4.8.268 sc

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **sc**

Stochastic Collocation as an emulator model.
Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Automated Refinement Type (Group 1)</td>
<td>p_refinement</td>
<td>Automatic polynomial order refinement</td>
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<tr>
<td>Optional</td>
<td></td>
<td>h_refinement</td>
<td>Employ h-refinement to refine the grid</td>
</tr>
<tr>
<td></td>
<td></td>
<td>max_refinement_iterations</td>
<td>Maximum number of expansion refinement iterations</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Interpolation Grid Type (Group 2)</td>
<td>quadrature_order</td>
<td>Order for tensor-products of Gaussian quadrature rules</td>
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<tr>
<td></td>
<td></td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
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<tr>
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<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>wiener</td>
<td>Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.</td>
</tr>
</tbody>
</table>
Optional

Covariance Type (Group 4)

diagonal_covariance

full_covariance

Use derivative data to construct surrogate models

Display only the diagonal terms of the covariance matrix

Display the full covariance matrix

Description

Selects stochastic collocation (SC) model to use in the Bayesian likelihood calculations. Most specification options are carried over for using SC as a surrogate within the Bayesian framework.

See Also

These keywords may also be of interest:

• stoch_collocation

Automatic polynomial order refinement

Specification

Alias: none

Argument(s): none

Default: no refinement

Child Keywords:
7.2 method

| dimension_adaptive | Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’. |

Description

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (`collocation_points` or `collocation_ratio`).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize `sobol`, `decay`, or `generalized` refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (`sobol` case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (`decay` case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized `dimension_adaptive` case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.4.8.270 uniform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- p_refinement
- uniform

Refine an expansion uniformly in all dimensions.
Specification

Alias: none
Argument(s): none

Description

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.4.8.271 dimension_adaptive

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- p_refinement
- dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Dimension Adaptivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>sobol Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher ‘importance’ in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.4.8.272 sobol

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- p_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol’ sensitivity indices.

Specification

Alias: none
Argument(s): none
Default: generalized

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.
Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

**Examples**

```plaintext
method, polynomial_chaos
    sparse_grid_level = 3
dimension_adaptive p_refinement generalized
    max_iterations = 20
    convergence_tol = 1.e-4
```

**h_refinement**

- **Keywords Area**
- **method**
- **bayes_calibration**
Employ h-refinement to refine the grid

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td><strong>Required (Choose One)</strong></td>
<td><strong>h-refinement Type</strong> (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

**Description**

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_-adaptive options.

7.2.68.4.8.275 uniform

- Keywords Area
- method
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias**: none  
**Argument(s)**: none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

**Dimension Adaptive**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias**: none  
**Argument(s)**: none

**Child Keywords:**

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tr>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

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Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher 'importance' in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.6.8.4.277 sobol

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- h_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol’ sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement sobol
    max_iterations = 20
    convergence_tol = 1.e-4
```

7.2.68.4.8.278  generalized

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- h_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

**Alias:** none

**Argument(s):** none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User’s Manual PCE descriptions for additional specifics).

Examples

```plaintext
method,
    polynomial_chaos
    sparse_grid_level = 3
    dimension_adaptive p_refinement generalized
    max_iterations = 20
    convergence_tol = 1.e-4
```
Planned future capability for local pointwise refinement within a generalized sparse grid.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

method,
    polynomial_chaos
dimension_adaptive generalized
p_refinement
    max_refinement_iterations = 20
    convergence_tol = 1.e-4
    sparse_grid_level = 1

7.2.68.4.8.281 quadrature_order

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• sc
• quadrature_order

Order for tensor-products of Gaussian quadrature rules

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td></td>
<td>Quadrature Rule Nesting (Group 1)</td>
<td>non_nested</td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

Description

Multidimensional integration by a tensor-product of Gaussian quadrature rules (specified with quadrature_order, and, optionally, dimension_preference). The default rule selection is to employ non_nested Gauss rules including Gauss-Hermite (for normals or transformed normals), Gauss-Legendre (for uniforms or transformed uniforms), Gauss-Jacobi (for betas), Gauss-Laguerre (for exponentials), generalized Gauss-Laguerre (for gammas), and numerically-generated Gauss rules (for other distributions when using an Extended basis). For the case of p_refinement or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse. By specifying a dimension_preference,
where higher preference leads to higher order polynomial resolution, the tensor grid may be rendered anisotropic. The dimension specified to have highest preference will be set to the specified `quadrature_order` and all other dimensions will be reduced in proportion to their reduced preference; any non-integral portion is truncated. To synchronize with tensor-product integration, a tensor-product expansion is used, where the order \( p_i \) of the expansion in each dimension is selected to be half of the integrand precision available from the rule in use, rounded down. In the case of non-nested Gauss rules with integrand precision \( 2m_i - 1 \), \( p_i \) is one less than the quadrature order \( m_i \) in each dimension (a one-dimensional expansion contains the same number of terms, \( p + 1 \), as the number of Gauss points). The total number of terms, \( N \), in a tensor-product expansion involving \( n \) uncertain input variables is

\[
N = 1 + P = \prod_{i=1}^{n} (p_i + 1)
\]

In some advanced use cases (e.g., multifidelity UQ), multiple grid resolutions can be employed; for this reason, the `quadrature_order` specification supports an array input.

A corresponding sequence specification is documented at, e.g., `quadrature_order_sequence` and `quadrature_order_sequence`.

**See Also**

These keywords may also be of interest:

- `quadrature_order_sequence` and `method-multifidelity_stoch_collocation-quadrature_order_sequence`

---

### 7.2.68.4.8.282 dimension_preference

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **sc**
- **quadrature_order**
- **dimension_preference**

A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.283 nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- quadrature_order
- nested

Enforce use of nested quadrature rules if available

Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.284 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- quadrature_order
- non_nested

Enforce use of non-nested quadrature rules
Specification

**Alias**: none

**Argument(s)**: none

**Description**

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.4.8.285 **sparse_grid_level**

- **Keywords** Area
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **sc**
- **sparse_grid_level**

Level to use in sparse grid integration or interpolation

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Group 1</td>
<td>nodal</td>
<td>Employ a nodal sparse grid construction in stochastic collocation</td>
</tr>
</tbody>
</table>
### Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with sparse_grid_level and, optionally, dimension_preference). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit non_nested specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the dimension_preference specification enables the use of anisotropic sparse grids (refer to the PCE description in the User’s Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full sparse_grid_level and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User’s Manual for additional details). As for quadrature_order, the sparse_grid_level specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., sparse_grid_level_sequence and sparse_grid_level_sequence

### See Also

These keywords may also be of interest:

- sparse_grid_level_sequence method-multifidelity_stoch_collocation-sparse_grid_level_sequence

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A set of weights specifying the relative importance of each uncertain variable (dimension)

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

**Description**

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

**See Also**

These keywords may also be of interest:

- sobol
- decay

Employ a nodal sparse grid construction in stochastic collocation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** nodal
Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.

7.2.68.4.8.288 hierarchical

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- sparse_grid_level
- hierarchical

Employ a hierarchical sparse grid construction

Specification

Alias: none
Argument(s): none

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

7.2.68.4.8.289 restricted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- sparse_grid_level
- restricted

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.
7.2 method

Specification

Alias: none

Argument(s): none

Default: restricted (except for generalized sparse grids)

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of $m = 2l + 1$ are used to enforce odd quadrature orders, where $l$ is the grid level and $m$ is the number of points in the rule. The precision of this Gauss rule is then $i = 2m - 1 = 4l + 1$. For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level $l$ for a sparse grid ($i = 4l + 1$) or an order $m$ for a tensor grid ($i = 2m - 1$). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.290 unrestricted

- Keywords Area
  - method
  - bayes_calibration
  - dream
  - emulator
  - sc
  - sparse_grid_level
  - unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none

Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of $m = 2l + 1$ are used to enforce odd quadrature orders, where $l$ is the grid level and $m$ is the number of points in the rule. The precision of this Gauss rule is then $i = 2m - 1 = 4l + 1$. For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level $l$ for a sparse grid ($i = 4l + 1$) or an order $m$ for a tensor grid ($i = 2m - 1$). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used: it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
sequences. To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension Adaptive p refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.291 nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- sparse_grid_level
- nested

Enforce use of nested quadrature rules if available

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.292 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- sparse_grid_level
- non_nested

Enforce use of non-nested quadrature rules

**Specification**

**Alias**: none

**Argument(s)**: none
Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.4.8.293  piecewise

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• sc
• piecewise

Use piecewise local basis functions

Specification

Alias: none

Argument(s): none

Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.4.8.294  askey

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• sc
• askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.4.8.296  use_derivatives

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.4.8.297  diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- sc
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
Specification

**Alias:** none

**Argument(s):** none

**Default:** diagonal\_covariance for response vector > 10; else full\_covariance

**Description**

With a large number of responses, the covariance matrix can be very large. \texttt{diagonal\_covariance} is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.4.8.298  \texttt{full\_covariance}

- Keywords Area
- method
- bayes\_calibration
- dream
- emulator
- sc
- full\_covariance

Display the full covariance matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

With a large number of responses, the covariance matrix can be very large. \texttt{full\_covariance} is used to force Dakota to output the full covariance matrix.

7.2.68.4.8.299  \texttt{mf\_sc}

- Keywords Area
- method
- bayes\_calibration
- dream
- emulator
- mf\_sc

Multifidelity Stochastic Collocation as an emulator model.
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
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<th>Required/Optional</th>
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<td>Automatic polynomial order refinement</td>
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<td>h_refinement</td>
<td></td>
<td>Employ h-refinement to refine the grid</td>
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<td>Optional</td>
<td>max_refinement_-iterations</td>
<td></td>
<td>Maximum number of expansion refinement iterations</td>
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<tr>
<td>Optional</td>
<td>allocation_control</td>
<td></td>
<td>Sample allocation approach for multifidelity expansions</td>
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<td>Optional</td>
<td>discrepancy_-emulation</td>
<td></td>
<td>Formulation for emulation of model discrepancies</td>
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<td>Required (Choose One)</td>
<td>Interpolation Grid Type (Group 2)</td>
<td>quadrature_order_-sequence</td>
<td>Sequence of quadrature orders used in a multi-stage expansion</td>
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<td></td>
<td>sparse_grid_level_-sequence</td>
<td>Sequence of sparse grid levels used in a multi-stage expansion</td>
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<td>Optional (Choose One)</td>
<td>Basis Polynomial Family (Group 3)</td>
<td>piecewise</td>
<td>Use piecewise local basis functions</td>
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<td></td>
<td>askey</td>
<td>Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.</td>
</tr>
</tbody>
</table>
7.2 method

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

<table>
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<tr>
<th>Optional</th>
<th>Covariance Type (Group 4)</th>
<th>use_derivatives</th>
<th>diagonal_covariance</th>
<th>full_covariance</th>
</tr>
</thead>
</table>

Optional (Choose One)

Description

Selects a multifidelity stochastic collocation (MF SC) surrogate model to use in the Bayesian likelihood calculations. Most specification options are carried over for using MF SC as a surrogate within the Bayesian framework.

See Also

These keywords may also be of interest:

- multifidelity_stoch_collocation

7.2.68.4.8.300  p_refinement

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- p_refinement

Specification

Alias: none

Argument(s): none

Default: no refinement

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>p-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.</td>
</tr>
</tbody>
</table>

**Description**

The `p_refinement` keyword specifies the usage of automated polynomial order refinement, which can be either `uniform` or `dimension_adaptive`.

The `dimension_adaptive` option is supported for the tensor-product quadrature and Smolyak sparse grid options and `uniform` is supported for tensor and sparse grids as well as regression approaches (collocation_points or collocation_ratio).

Each of these refinement cases makes use of the `max_iterations` and `convergence_tolerance` method independent controls. The former control limits the number of refinement iterations, and the latter control terminates refinement when the two-norm of the change in the response covariance matrix (or, in goal-oriented approaches, the two-norm of change in the statistical quantities of interest (QOI)) falls below the tolerance.

The `dimension_adaptive` case can be further specified to utilize sobol, decay, or generalized refinement controls. The former two cases employ anisotropic tensor/sparse grids in which the anisotropic dimension preference (leading to anisotropic integrations/expansions with differing refinement levels for different random dimensions) is determined using either total Sobol' indices from variance-based decomposition (sobol case: high indices result in high dimension preference) or using spectral coefficient decay rates from a rate estimation technique similar to Richardson extrapolation (decay case: low decay rates result in high dimension preference). In these two cases as well as the `uniform` refinement case, the `quadrature_order` or `sparse_grid_level` are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the `uniform` refinement case with regression approaches, the `expansion_order` is ramped by one on each iteration while the oversampling ratio (either defined by `collocation_ratio` or inferred from `collocation_points` based on the initial expansion) is held fixed. Finally, the generalized dimension_adaptive case is the default adaptive approach; it refers to the generalized sparse grid algorithm, a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

For the case of `p_refinement` or the case of an explicit nested override, Gauss-Hermite rules are replaced with Genz-Keister nested rules and Gauss-Legendre rules are replaced with Gauss-Patterson nested rules, both of which exchange lower integrand precision for greater point reuse.

7.2.68.4.8.301 uniform

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
• mf_sc

• p_refinement

• uniform

Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

7.2.68.4.8.302 dimension_adaptive

• Keywords Area

• method

• bayes_calibration

• dream

• emulator

• mf_sc

• p_refinement

• dimension_adaptive

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher 'importance' in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.4.8.303 sobol

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_sc**
- **p_refinement**
- **dimension_adaptive**
- **sobol**

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

### Specification

**Alias:** none

**Argument(s):** none

**Default:** generalized

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Dimension Adaptivity Estimation Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>sobol</td>
<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>
7.2 method

Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.4.8.304 generalized

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- p_refinement
- dimension_adaptive
- generalized

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none
Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

Examples

method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
Employ h-refinement to refine the grid

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no refinement  
**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td>h-refinement Type (Group 1)</td>
<td>uniform</td>
<td>Refine an expansion uniformly in all dimensions.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dimension_-adaptive</td>
<td>Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher ‘importance’.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_adaptive</td>
<td>Planned future capability for local pointwise refinement within a generalized sparse grid.</td>
</tr>
</tbody>
</table>

**Description**

Automated expansion refinement can be selected as either p_refinement or h_refinement, and either refinement specification can be either uniform or dimension_adaptive. The dimension_adaptive case can be further specified as either sobol or generalized (decay not supported). Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls. The h_refinement specification involves use of the same piecewise interpolants (linear or cubic Hermite splines) described above for the piecewise specification option (it is not necessary to redundantly specify piecewise in the case of h_refinement). In future releases, the hierarchical interpolation approach will enable local refinement in addition to the current uniform and dimension_-adaptive options.
Refine an expansion uniformly in all dimensions.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The quadrature_order or sparse_grid_level are ramped by one on each refinement iteration until either of the two convergence controls is satisfied. For the uniform refinement case with regression approaches, the expansion_order is ramped by one on each iteration while the oversampling ratio (either defined by collocation_ratio or inferred from collocation_points based on the initial expansion) is held fixed.

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to have higher 'importance'.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
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<tr>
<td>Required (Choose One)</td>
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<td>Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>generalized</td>
<td>Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.</td>
</tr>
</tbody>
</table>

**Description**

Perform anisotropic expansion refinement by preferentially adapting in dimensions that are detected to hold higher 'importance' in resolving statistical quantities of interest.

Dimension importance must be estimated as part of the refinement process. Techniques include either sobol or generalized for stochastic collocation and either sobol, decay, or generalized for polynomial chaos. Each of these automated refinement approaches makes use of the max_iterations and convergence_tolerance iteration controls.

7.2.68.4.8.308 sobol

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- h_refinement
- dimension_adaptive
- sobol

Estimate dimension preference for automated refinement of stochastic expansion using total Sobol' sensitivity indices.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** generalized
Description

Determine dimension preference for refinement of a stochastic expansion from the total Sobol' sensitivity indices obtained from global sensitivity analysis. High indices indicate high importance for resolving statistical quantities of interest and therefore result in high dimension preference.

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement sobol
  max_iterations = 20
  convergence_tol = 1.e-4

7.2.68.4.8.309 generalized

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_sc
• h_refinement
• dimension_adaptive
• generalized
```

Use the generalized sparse grid dimension adaptive algorithm to refine a sparse grid approximation of stochastic expansion.

Specification

Alias: none

Argument(s): none

Description

The generalized sparse grid algorithm is a greedy approach in which candidate index sets are evaluated for their impact on the statistical QOI, the most influential sets are selected and used to generate additional candidates, and the index set frontier of a sparse grid is evolved in an unstructured and goal-oriented manner (refer to User's Manual PCE descriptions for additional specifics).

Examples

```plaintext
method, polynomial_chaos
  sparse_grid_level = 3
  dimension_adaptive p_refinement generalized
  max_iterations = 20
  convergence_tol = 1.e-4
```
Planned future capability for local pointwise refinement within a generalized sparse grid.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Sparse grid interpolation admits approaches for pointwise local refinement within the general framework of generalized sparse grids. This algorithmic capability is currently in a partial prototype stage.

Maximum number of expansion refinement iterations

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

**Description**

Limits the number of times the expansion can be refined under various refinement strategies.
Examples

method,  
   polynomial_chaos  
      dimension_adaptive generalized  
      p_refinement  
         max_refinement_iterations = 20  
         convergence_tol = 1.e-4  
         sparse_grid_level = 1

7.2.6.8.4.8.312 allocation_control

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_sc**
- **allocation_control**

Sample allocation approach for multifidelity expansions

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**: Required/Optional | Description of Group | Dakota Keyword | Dakota Keyword Description  
---|---|---|---
**Required**(Choose One) | Multifidelity Sample Allocation Control (Group 1) | greedy | Sample allocation based on greedy refinement within multifidelity stochastic collocation

Description

Multifidelity surrogate approaches, including polynomial chaos, stochastic collocation, and function train, can optionally employ a integrated greedy competition across the model sequence, where each model index can supply one or more refinement candidates which are competed to determine the candidate with the greatest impact on the QoI statistics per unit cost. This greedy competition implicitly determines the optimal sample allocation across model indices.

**Default Behavior**

The default, when allocation_control is not specified, is to compute or adapt separately for each model index (individual instead of integrated refinement).
Multifidelity stochastic collocation supports greedy refinement strategies using tensor and sparse grids for both nodal and hierarchical collocation approaches. The key idea is that each level of the model hierarchy being approximated can generate one or more candidates for refinement. These candidates are competed against each other within an integrated competition, and the candidate that induces the largest change in the statistical QoI (response covariance by default, or results of any level mappings when specified), normalized by relative cost of evaluating the candidate, is selected and then used to generate additional candidates for consideration at its model level.

The following example of greedy multifidelity stochastic collocation using nodal interpolation starts from a zeroth-order expansion (a constant) for each level, and generates uniform candidate refinements for each level that are competed in a greedy competition. The number of new samples for the incremented candidate expansion order is determined from the quadrature rules of the new sparse grid level. In this case, the number of candidates for each level is limited to one uniform refinement of the current sparse grid level.

```dakota
method,
  model_pointer = ‘HIERARCH’
  multifidelity_stoch_collocation
    nodal
    allocation_control greedy
    p_refinement uniform
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-3
```

The next example employs generalized sparse grids and hierarchical interpolation. Each level starts from a level 0 reference grid (a single point) and generates multiple admissible index set candidates. The full set of candidates across all levels is competed within a unified greedy competition, where the greedy selection metric is the induced change in the statistical QoI, normalized by the aggregate simulation cost of the index set candidate. In this case, there are multiple candidates for each level and the number of candidates grows rapidly with random dimension and grid level.

```dakota
method,
  model_pointer = ‘HIERARCH’
  multifidelity_stoch_collocation
    hierarchical
    allocation_control greedy
    p_refinement uniform
    sparse_grid_level_sequence = 0 unrestricted
    convergence_tolerance 1.e-3
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Formulation for emulation of model discrepancies.

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** distinct  
**Child Keywords:**

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<td>Distinct formulation for emulation of model discrepancies.</td>
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<tr>
<td></td>
<td></td>
<td>recursive</td>
<td>Recursive formulation for emulation of model discrepancies.</td>
</tr>
</tbody>
</table>

**Description**

In many uncertainty quantification approaches, model discrepancies are emulated using, e.g., polynomial chaos, stochastic collocation, or Gaussian process models. Two formulations are available for this emulation:

1. **distinct** emulation (default), in which we directly approximate the difference or ratio between the evaluations of two models or solution levels.
2. **recursive** emulation (experimental option), in which we approximate a difference or ratio among the new model evaluation and the emulator approximation of the previous model.
The latter approach is a form of hierarchical emulation in which we emulate the surplus between the previous emulator and the new modeling level. This approach has a few advantages: (i) it reduces bias by correcting for emulation errors that occur at previous levels, and (ii) it does not require paired model evaluations for each discrepancy level, which reduces cost, allows for disparate sample points, and simplifies data imports.

On the other hand, its primary disadvantage is that the aggregate emulation is only as good as its weakest link, in that a poor emulator recovery can create difficulty in accurately resolving discrepancies that are recursively dependent on it. Thus, the distinct approach may tend to be more expensive in exchange for greater robustness.

**Examples**

```plaintext
method,
    multilevel_polynomial_chaos
    expansion_order_sequence = 2
    collocation_ratio = .9
    orthogonal_matching_pursuit
    discrepancy_emulation recursive
```

**Distinct**

Distinct formulation for emulation of model discrepancies.

**Specification**

- **Alias:** paired
- **Argument(s):** none

**Description**

The distinct approach to discrepancy emulation directly approximate the difference or ratio between the evaluations of two models or solution levels. Refer to the parent documentation node for additional discussion.

**Examples**

```plaintext
7.2.68.4.8.315  recursive
    • Keywords Area
    • method
    • bayes_calibration
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Recursive formulation for emulation of model discrepancies.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The recursive approach to discrepancy emulation approximates a difference or ratio among the new model evaluation and the emulator approximation of the previous model. Refer to the parent documentation node for additional discussion.

**Examples**

7.2.68.4.8.317 quadrature_order_sequence

- **Keywords Area**
- **method**
- **bayes_calibration**
- **dream**
- **emulator**
- **mf_sc**
- **quadrature_order_sequence**

Sequence of quadrature orders used in a multi-stage expansion

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Keywords Area

<table>
<thead>
<tr>
<th>Optional</th>
<th>Quadrature Rule Nesting (Group 1)</th>
<th>A set of weights specifying the relative importance of each uncertain variable (dimension)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dimension_preference</td>
<td>nested</td>
<td>Enforce use of nested quadrature rules if available</td>
</tr>
<tr>
<td>non_nested</td>
<td></td>
<td>Enforce use of non-nested quadrature rules</td>
</tr>
</tbody>
</table>

### Description

Each level entry of the `quadrature_order_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support quadrature order sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `quadrature_order`

### See Also

These keywords may also be of interest:

- `quadrature_order`

### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids
Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

- sobol
- decay

7.2.68.4.8.319 nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- quadrature_order_sequence
- nested

Enforce use of nested quadrature rules if available

Specification

**Alias:** none

**Argument(s):** none

**Default:** quadrature: non_nested unless automated refinement; sparse grids: nested

Description

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

7.2.68.4.8.320 non_nested

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- quadrature_order_sequence
- non_nested

Enforce use of non-nested quadrature rules
Specification

Alias: none
Argument(s): none

Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule

7.2.68.4.8.321 sparse_grid_level_sequence

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_sc
• sparse_grid_level_sequence

Sequence of sparse grid levels used in a multi-stage expansion

Specification

Alias: none
Argument(s): INTEGERLIST

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>nodal</td>
<td>Employ a nodal sparse grid construction in stochastic collocation</td>
</tr>
</tbody>
</table>
Each level entry of the `sparse_grid_level_sequence` applies to one expansion within a multi-stage expansion. Current multi-stage expansions that support sparse grid level sequences include multilevel polynomial chaos, multifidelity polynomial chaos, or multifidelity stochastic collocation expansions.

If adaptive refinement is active, then this sequence specifies the starting point for each level within either an individual or integrated refinement approach.

A corresponding scalar specification is documented at, e.g., `sparse_grid_level`
Specification

Alias: none
Argument(s): REALLIST
Default: isotropic grids

Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.

See Also

These keywords may also be of interest:

• sobol
• decay

7.2.68.4.8.323 nodal

• Keywords Area
• method
• bayes_calibration
• dream
• emulator
• mf_sc
• sparse_grid_level_sequence
• nodal

Employ a nodal sparse grid construction in stochastic collocation

Specification

Alias: none
Argument(s): none
Default: nodal

Description

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The nodal formulation does not require nested interpolation rules and is therefore more generally applicable. However, in the context of adaptive refinement, it can suffer from loss of precision due to subtractive cancellation.
Employ a hierarchical sparse grid construction

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Stochastic collocation with sparse grids can employ either a nodal or hierarchical formulation. The hierarchical formulation requires nested interpolation rules as it utilizes increments in collocation points along with hierarchical weights defined from a hierarchical basis formulation.

This keyword requests the hierarchical formulation, which has benefits for adaptive refinement by supporting direct computation of increments in targeted statistics (e.g. mean, covariance, reliability indices), avoiding loss of precision from subtractive cancellation.

Restrict the growth rates for nested and non-nested rules can be synchronized for consistency.

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** restricted (except for generalized sparse grids)
Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.

7.2.68.4.8.326 unrestricted

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- sparse_grid_level_sequence
- unrestricted

Override the default restriction of growth rates for nested and non-nested rules that are by default synchronized for consistency.

Specification

Alias: none
Argument(s): none

Description

In the quadrature and sparse grid cases, growth rates for nested and non-nested rules can be synchronized for consistency. For a non-nested Gauss rule used within a sparse grid, linear one-dimensional growth rules of \( m = 2l + 1 \) are used to enforce odd quadrature orders, where \( l \) is the grid level and \( m \) is the number of points in the rule. The precision of this Gauss rule is then \( i = 2m - 1 = 4l + 1 \). For nested rules, order growth with level is typically exponential; however, the default behavior is to restrict the number of points to be the lowest order rule that is available that meets the one-dimensional precision requirement implied by either a level \( l \) for a sparse grid (\( i = 4l + 1 \)) or an order \( m \) for a tensor grid (\( i = 2m - 1 \)). This behavior is known as "restricted growth" or "delayed sequences." To override this default behavior in the case of sparse grids, the unrestricted keyword can be used; it cannot be overridden for tensor grids using nested rules since it also provides a mapping to the available nested rule quadrature orders. An exception to the default usage of restricted growth is the dimension_adaptive p_refinement generalized sparse grid case described previously, since the ability to evolve the index sets of a sparse grid in an unstructured manner eliminates the motivation for restricting the exponential growth of nested rules.
Enforce use of nested quadrature rules if available

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Default**: quadrature: non_nested unless automated refinement; sparse grids: nested

**Description**

Enforce use of nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the Nested Genz-Keister rule instead of the default non-nested Gauss-Hermite rule variables are

Enforce use of non-nested quadrature rules

**Specification**

- **Alias**: none
- **Argument(s)**: none
Description

Enforce use of non-nested quadrature rules if available. For instance if the aleatory variables are Gaussian use the non-nested Gauss-Hermite rule.

7.2.68.4.8.329  piecewise

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- piecewise

Use piecewise local basis functions.

Specification

Alias: none
Argument(s): none
Default: extended (Askey + numerically-generated)

Description

SC also supports the option of piecewise local basis functions. These are piecewise linear splines, or in the case of gradient-enhanced interpolation via the use_derivatives specification, piecewise cubic Hermite splines. Both of these basis selections provide local support only over the range from the interpolated point to its nearest 1D neighbors (within a tensor grid or within each of the tensor grids underlying a sparse grid), which exchanges the fast convergence of global bases for smooth functions for robustness in the representation of nonsmooth response functions (that can induce Gibbs oscillations when using high-order global basis functions). When local basis functions are used, the usage of nonequidistant collocation points (e.g., the Gauss point selections described above) is not well motivated, so equidistant Newton-Cotes points are employed in this case, and all random variable types are transformed to standard uniform probability space.

7.2.68.4.8.330  askey

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- askey

Select the standardized random variables (and associated basis polynomials) from the Askey family that best match the user-specified random variables.
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Askey option employs standard normal, standard uniform, standard exponential, standard beta, and standard gamma random variables in a transformed probability space. These selections correspond to Hermite, Legendre, Laguerre, Jacobi, and generalized Laguerre orthogonal polynomials, respectively.

Specific mappings for the basis polynomials are based on a closest match criterion, and include Hermite for normal (optimal) as well as bounded normal, lognormal, bounded lognormal, gumbel, frechet, and weibull (sub-optimal); Legendre for uniform (optimal) as well as loguniform, triangular, and bin-based histogram (sub-optimal); Laguerre for exponential (optimal); Jacobi for beta (optimal); and generalized Laguerre for gamma (optimal).

See Also

These keywords may also be of interest:

- polynomial_chaos
- wiener

Use standard normal random variables (along with Hermite orthogonal basis polynomials) when transforming to a standardized probability space.

Specification

**Alias:** none

**Argument(s):** none

**Default:** extended (Askey + numerically-generated)

Description

The Wiener option employs standard normal random variables in a transformed probability space, corresponding to a Hermite orthogonal polynomial basis. This is the same nonlinear variable transformation used by local and global reliability methods (and therefore has the same variable support).
See Also

These keywords may also be of interest:

- polynomial_chaos
- askey

7.2.68.4.8.332  use_derivatives

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- use_derivatives

Use derivative data to construct surrogate models

Specification

**Alias:** none

**Argument(s):** none

**Default:** use function values only

Description

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, its use with Surfpack Gaussian process is not recommended.

7.2.68.4.8.333  diagonal_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- diagonal_covariance

Display only the diagonal terms of the covariance matrix
7.2 method

Specification

Alias: none

Argument(s): none

Default: diagonal_covariance for response vector > 10; else full_covariance

Description

With a large number of responses, the covariance matrix can be very large. diagonal_covariance is used to suppress the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

7.2.68.4.8.3 full_covariance

- Keywords Area
- method
- bayes_calibration
- dream
- emulator
- mf_sc
- full_covariance

Display the full covariance matrix

Specification

Alias: none

Argument(s): none

Description

With a large number of responses, the covariance matrix can be very large. full_covariance is used to force Dakota to output the full covariance matrix.

7.2.68.4.9 standardized_space

- Keywords Area
- method
- bayes_calibration
- dream
- standardized_space

Perform Bayesian inference in standardized probability space

Specification

Alias: none

Argument(s): none
Description

This option transforms the inference process (MCMC sampling and any emulator model management) into a standardized probability space.

The variable transformations performed are as described in askey.

Default Behavior

The default for the Gaussian process and no emulator options is to perform inference in the original probability space (no transformation). Polynomial chaos and stochastic collocation emulators, on the other hand, are always formed in standardized probability space, such that the inference process is also performed in this standardized space.

Expected Output

The user will see the truth model evaluations performed in the original space, whereas any method diagnostics relating to the MCMC samples (e.g., QUESO data in the outputData directory) will report points and response data (response gradients and Hessians, if present, will differ but response values will not) that correspond to the transformed space.

Usage Tips

Selecting standardized_space generally has the effect of scaling the random variables to be of comparable magnitude, which can improve the efficiency of the Bayesian inference process.

Examples

```
method, bayes_calibration queso
  samples = 2000 seed = 348
  dram
  standardized_space
```

7.2.68.4.10 export_chain_points_file

- Keywords Area
- method
- bayes_calibration
- dream
- export_chain_points_file

Export the MCMC chain to the specified filename

Specification

Alias: none

Argument(s): STRING

Default: chain export to default filename

Child Keywords:
7.2 method

<table>
<thead>
<tr>
<th>Optional (Choose One)</th>
<th>Tabular Format (Group 1)</th>
<th>custom_annotated</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

The filename to which the final MCMC posterior chain will be exported.

**Default Behavior**  No export to file.

**Expected Output**

A tabular data file will be produced in the specified format (annotated by default) containing samples from the posterior distribution.

**Usage Tips**

**Additional Discussion**

7.2.68.4.10.1 custom_annotated

- Keywords Area
  - method
  - bayes_calibration
  - dream
  - export_chain_points_file
  - custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
    tabular_data
        tabular_data_file = 'dakota_summary.dat'
        custom_annotated header eval_id

Resulting tabular file:

%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1    0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.68.4.10.2 header

- Keywords Area
- method
7.2 method

- bayes_calibration
- dream
- export_chain_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.4.10.3  eval_id

- Keywords Area
- method
- bayes_calibration
- dream
- export_chain_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.4.10.4  interface_id

- Keywords Area
- method
- bayes_calibration
- dream
- export_chain_points_file
Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

See description of parent `custom_annotated`

7.2.68.4.10.5  annotated

- Keywords Area
- method
- bayes_calibration
- dream
- export_chain_points_file
- annotated

Selects annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**
• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.4.10.6 freeform

• Keywords Area
  • method
  • bayes_calibration
  • dream
  • export_chain_points_file
  • freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated
Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
```

7.2.68.5  muq

- Keywords Area
- method
- bayes_calibration
- muq

Markov Chain Monte Carlo algorithms from the MUQ package

Topics

This keyword is related to the topics:

- bayesian_calibration
7.2 method

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>chain_samples</td>
<td></td>
<td>Number of Markov Chain Monte Carlo posterior samples</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td></td>
<td>Selection of a random number generator</td>
</tr>
</tbody>
</table>

Optional (Choose One) MCMC Algorithm (Group 1)

- dram: Use the DRAM MCMC algorithm
- delayed_rejection: Use the Delayed Rejection MCMC algorithm
- adaptive_-metropolis: Use the Adaptive Metropolis MCMC algorithm
- metropolis_-hastings: Use the Metropolis-Hastings MCMC algorithm

Optional

- proposal_-covariance: Defines the technique used to generate the MCMC proposal covariance.

Description

The muq method supports the following MCMC algorithms: adaptive metropolis (AM), Metropolis Hasting (MH), delayed rejection (DR), or delayed-rejection adaptive metropolis (DRAM).

The muq method is currently an experimental method that relies on algorithms from MIT’s MUQ code documented at: https://bitbucket.org/mituq/muq2/src/master/

We anticipate using more advanced features of MUQ such as Hamiltonian Monte Carlo and Langevin methods in future releases of Dakota.

7.2.68.5.1 chain_samples

- Keywords Area
- method
- bayes_calibration
Number of Markov Chain Monte Carlo posterior samples

**Specification**

**Alias:** samples  
**Argument(s):** INTEGER  
**Default:** method-dependent

**Description**

The `chain_samples` keyword indicates the number of draws from the posterior distribution to perform. When an emulator is active, this will be the number of samples on the constructed surrogate model.

**Default Behavior**

The default number of chain samples is method-dependent. QUESO and MUQ methods use 1000. DREAM uses (number of generations) x (number of chains), resulting in `chain_samples` close to that specified.

**Usage Tips**

MCMC methods typically require a large number of chain samples to converge, often thousands to millions.

**Examples**

```
method bayes_calibration queso
    chain_samples = 20000
```

---

**7.2.68.5.2 seed**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **muq**
- **seed**

Seed of the random number generator

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** system-generated (non-repeatable)
Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output

If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

Usage Tips

If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.6.8.5.3 rng

- Keywords Area
  - method
  - bayes_calibration
  - muq
  - rng

Selection of a random number generator

Specification

Alias: none

Argument(s): none

Default: Mersenne twister ( mt19937 )

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
<td></td>
</tr>
</tbody>
</table>
Description

The `rng` keyword is used to indicate a choice of random number generator.

**Default Behavior**

If specified, the `rng` keyword must be accompanied by either `rnum2` (pseudo-random numbers) or `mt19937` (random numbers generated by the Mersenne twister). Otherwise, `mt19937`, the Mersenne twister is used by default.

**Usage Tips**

The default is recommended, as the Mersenne twister is a higher quality random number generator.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
7.2.68.5.3.1 mt19937
  • Keywords Area
  • method
  • bayes_calibration
  • muq
  • rng
  • mt19937
Generates random numbers using the Mersenne twister
```

**Specification**

- **Alias**: none
- **Argument(s)**: none

**Description**

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

**Default Behavior**

`mt19937` is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

**Usage Tips**

Use of the Mersenne twister random number generator (`mt19937`) is recommended.
Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.68.5.3.2 rnum2

- Keywords Area
- method
- bayes_calibration
- muq
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

**Alias:** none

**Argument(s):** none

Description

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.68.5.4 dram

- Keywords Area
- method
- bayes_calibration
- muq
- dram

Use the DRAM MCMC algorithm
Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none
Argument(s): none
Default: dram

Description

The type of Markov Chain Monte Carlo used. This keyword specifies the use of DRAM, (Delayed Rejection Adaptive Metropolis)[40].

Default Behavior

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Usage Tips

If the user knows very little about the proposal covariance, using dram is a recommended strategy. The proposal covariance is adaptively updated, and the delayed rejection may help improve low acceptance rates.

Examples

```
method, bayes_calibration queso
dram
samples = 10000 seed = 348
```

7.2.68.5.5 delayed_rejection

- Keywords Area
- method
- bayes_calibration
- muq
- delayed_rejection

Use the Delayed Rejection MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** dram

**Description**

This keyword specifies the use of the Delayed Rejection algorithm in which there can be a delay in rejecting samples from the chain. That is, the "DR" part of DRAM is used but the "AM" part is not, rather a regular Metropolis-Hastings algorithm is used.

**Default Behavior**

Five MCMC algorithm variants are supported: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

**Usage Tips**

If the user knows something about the proposal covariance or the proposal covariance is informed through derivative information, using delayed_rejection is preferred over dram: the proposal covariance is already being informed by derivative information and the adaptive metropolis is not necessary.

**Examples**

```
method,  
bayes_calibration queso  
delayed_rejection  
samples = 10000 seed = 348
```

**See Also**

These keywords may also be of interest:

- proposal_covariance

7.2.68.5.6 adaptive_metropolis

- Keywords Area
  - method
  - bayes_calibration
  - muq
  - adaptive_metropolis

Use the Adaptive Metropolis MCMC algorithm

**Topics**

This keyword is related to the topics:

- bayesian_calibration
Keywords Area

Specification

Alias: none
Argument(s): none
Default: dram

Description

This keyword specifies the use of the Adaptive Metropolis algorithm. That is, the "AM" part of DRAM is used but the "DR" part is not: specifying this keyword activates only the Adaptive Metropolis part of the MCMC algorithm, in which the covariance of the proposal density is updated adaptively.

Default Behavior

Five MCMC algorithm variants are supported in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Four MCMC variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings and adaptive_metropolis.

Usage Tips

If the user knows very little about the proposal covariance, but doesn’t want to incur the cost of using full dram with both delayed rejection and adaptive metropolis, specifying only adaptive_metropolis offers a good strategy.

Examples

```plaintext
method,
    bayes_calibration queso
    adaptive_metropolis
    samples = 10000 seed = 348
```

7.2.68.5.7 metropolis_hastings

- Keywords Area
- method
- bayes_calibration
- muq
- metropolis_hastings

Use the Metropolis-Hastings MCMC algorithm

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none
Argument(s): none
Default: dram
Description

This keyword specifies the use of a Metropolis-Hastings algorithm for the MCMC chain generation. This means there is no delayed rejection and no adaptive proposal covariance updating as in DRAM.

Default Behavior

Five MCMC algorithm variants are supported currently in QUESO: dram, delayed_rejection, adaptive_metropolis, metropolis_hastings, and multilevel. The default is dram.

Four MCMC algorithm variants are currently supported in MUQ: dram, delayed_rejection, metropolis_hastings and adaptive_metropolis.

Usage Tips

If the user wants to use Metropolis-Hastings, possibly as a comparison to the other methods which involve more chain adaptation, this is the MCMC type to use.

Examples

```
method,
    bayes_calibration queso
    metropolis_hastings
    samples = 10000 seed = 348
```

7.2.68.5.8 proposal_covariance

- Keywords Area

- method

- bayes_calibration

- muq

- proposal_covariance

Defines the technique used to generate the MCMC proposal covariance.

Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Proposal Covariance Source (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>prior</td>
<td>Uses the covariance of the prior distributions to define the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>derivatives</td>
<td>Use derivatives to inform the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>values</td>
<td>Specifies matrix values to use as the MCMC proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>filename</td>
<td>Uses a file to import a user-specified MCMC proposal covariance.</td>
</tr>
</tbody>
</table>

**Description**

The proposal covariance is used to define a multivariate normal (MVN) jumping distribution used to create new points within a Markov chain. That is, a new point in the chain is determined by sampling within a MVN probability density with prescribed covariance that is centered at the current chain point. The accuracy of the proposal covariance has a significant effect on rejection rates and the efficiency of chain mixing.

**Default Behavior**

The default proposal covariance is `prior` when no emulator is present; `derivatives` when an emulator is present.

**Expected Output**

The effect of the proposal covariance is reflected in the MCMC chain values and the rejection rates, which can be seen in the diagnostic outputs from the QUESO solver within the `QuesoDiagnostics` directory.

**Usage Tips**

When derivative information is available inexpensively (e.g., from an emulator model), the derived-based proposal covariance forms a more accurate proposal distribution, resulting in lower rejection rates and faster chain mixing.

7.2.68.5.8.1 prior

- Keywords Area
- method
- bayes_calibration
- muq
- proposal_covariance
- prior

Uses the covariance of the prior distributions to define the MCMC proposal covariance.
Topics

This keyword is related to the topics:

- bayesian_calibration

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>multiplier</td>
<td>Multiplier to scale prior variance</td>
</tr>
</tbody>
</table>

Description

This keyword selection results in definition of the MCMC proposal covariance from the covariance of the prior distributions. This covariance is currently assumed to be diagonal without correlation.

Default Behavior

This is the default proposal_covariance option.

Usage Tips

Since this proposal covariance is defined globally, the chain does not need to be periodically restarted using local updates to this proposal. However, it is usually effective to adapt the proposal using one of the adaptive metropolis MCMC options.

Examples

```
method,
    bayes_calibration queso
    samples = 2000 seed = 348
dram
proposal_covariance prior
```

7.2.68.5.8.2 multiplier

- Keywords Area
- method
- bayes_calibration
- muq
- proposal_covariance
- prior
- multiplier

Multiplier to scale prior variance
Specification

Alias: none

Argument(s): REAL

Default: 1.0

Description

The initial proposal covariance will be given by the prior variance times the multiplier.

Default Behavior  When using prior-based proposal covariance, the default is to use the prior variance for the proposal (multiplier = 1.0)

Usage Tips

The prior variance may result in too tight or narrow a proposal covariance. The multiplier can be used to scale all entries of the prior variance in determining the initial proposal covariance.

Examples

method bayes_calibration queso
    samples = 2000 seed = 348
dram
    proposal_covariance prior
        multiplier 0.1

Topics

This keyword is related to the topics:

• bayesian_calibration

Use derivatives to inform the MCMC proposal covariance.

Child Keywords: none
### Description

This keyword selection results in definition of the MCMC proposal covariance from the Hessian of the misfit function (negative log likelihood), where this Hessian is defined from either a Gauss-Newton approximation (using only first derivatives of the calibration terms) or a full Hessian (using values, first derivatives, and second derivatives of the calibration terms). If this Hessian is indeterminate, it will be corrected as described in [16].

**Default Behavior**  The default is prior based proposal covariance. This is a more advanced option that exploits structure in the form of the likelihood.

**Expected Output**

When derivatives are specified for defining the proposal covariance, the misfit Hessian and its inverse (the MVN proposal covariance) will be output to the standard output stream.

**Usage Tips**

The full Hessian of the misfit is used when either supported by the emulator in use (for PCE and surfpack GP, but not SC or dakota GP) or by the user’s response specification (Hessian type is not "no_hessians"), in the case of no emulator. When this full Hessian is indefinite and cannot be inverted to form the proposal covariance, fallback to the positive semi-definite Gauss-Newton Hessian is employed.

Since this proposal covariance is locally accurate, it should be updated periodically using the `update_period` option. While the adaptive metropolis option can be used in combination with derivative-based preconditioning, it is generally preferable to instead decrease the proposal update period due to the improved local accuracy of this approach.

**Examples**

Generate a 2000 sample posterior chain, using derivatives to initialize the proposal covariance at the start of the chain.

```
method,
    bayes_calibration queso
    samples = 2000 seed = 348
    delayed_rejection
    emulator pce sparse_grid_level = 2
    proposal_covariance derivatives

7.2.68.5.8.4 update_period
```

**Period at which to update derivative-based proposal covariance**
Specification

**Alias:** none

**Argument(s):** INTEGER

**Description**

For derivative-based proposal covariance, this specifies the period (number of accepted MCMC samples) after which the proposal covariance is updated using derivative values at the current chain point.

**Default Behavior**

When `update_period` is not specified, derivatives will inform the proposal covariance at the start of the chain, but not updated further.

**Usage Tips**

The `update_period` should be tailored to the size of the total chain, accounting for the relative expense of derivative-based proposal updates.

**Examples**

```
method,
    bayes_calibration queso
    samples = 2000 seed = 348
    delayed_rejection
    emulator pce sparse_grid_level = 2
    proposal_covariance derivatives
    update_period = 40  # update proposal covariance every 40 points
```

7.2.68.5.8.5 values

- **Keywords Area**

- **method**

- **bayes_calibration**

- **muq**

- **proposal_covariance**

- **values**

Specifies matrix values to use as the MCMC proposal covariance.

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**
### Description
This keyword selection results in definition of the MCMC proposal covariance from user-specified matrix values. The matrix input format must be declared as either a full matrix or a matrix diagonal.

### Default Behavior
This option is not the default, and generally implies special a priori knowledge from the user.

### Usage Tips
This option is not supported for the case of transformations to standardized probability space.

### Examples
```
method,
   bayes_calibration queso
   samples = 1000 seed = 348
   dram
   proposal_covariance
      values ... # See leaf nodes for required format option
```

#### 7.2.68.5.8.6 diagonal
- Keywords Area
- method
- bayes_calibration
- muq
- proposal_covariance
- values
- diagonal

Specifies the diagonal matrix format when specifying a user-specified proposal covariance.

### Specification
**Alias:** none
**Argument(s):** none
Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.

Examples

```plaintext
method,
  bayes_calibration queso
  samples = 1000 seed = 348
  dram
  proposal_covariance
    diagonal values 1.0e6 1.0e-1
```

7.2.68.5.8.7 matrix

- Keywords Area
- method
- bayes_calibration
- muq
- proposal_covariance
- values
- matrix

Specifies the full matrix format when specifying a user-specified proposal covariance.

Specification

**Alias**: none

**Argument(s)**: none

Description

When specifying the MCMC proposal covariance in an input file, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.

Examples

```plaintext
method,
  bayes_calibration queso
  samples = 1000 seed = 348
  dram
  proposal_covariance
    matrix values 1.0 0.1
                  0.1 2.0
```

7.2.68.5.8.8 filename

- Keywords Area
- method
- bayes_calibration
7.2 method

- muq
- proposal_covariance
- filename

Uses a file to import a user-specified MCMC proposal covariance.

Specification

Alias: none

Argument(s): STRING

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Values For (Group 1)</th>
<th>Dakota Keyword diagonal</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Specifies the diagonal matrix format when importing a user-specified proposal covariance.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>matrix</td>
<td>Specifies the full matrix format when importing a user-specified proposal covariance.</td>
</tr>
</tbody>
</table>

Description

This keyword selection results in definition of the MCMC proposal covariance from importing data a user-specified filename. This import must be declared as either a full matrix or a matrix diagonal.

Default Behavior

This option is not the default, and generally implies special a priori knowledge from the user.

Usage Tips

This option is not supported for the case of transformations to standardized probability space.

Examples

method,
bayes_calibration queso
samples = 1000 seed = 348
dram
proposal_covariance
filename ... # See leaf nodes for required format option

7.2.68.5.8.9 diagonal

- Keywords Area
- method
Specifies the diagonal matrix format when importing a user-specified proposal covariance.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a diagonal matrix format, i.e., the user only provides the (positive) values along the diagonal.

**Examples**

```plaintext
method,  
    bayes_calibration queso  
    samples = 1000 seed = 348  
    dram  
    proposal_covariance  
        diagonal filename 'dakota_cantilever_queso.diag.dat'
```

7.2.68.5.8.10  

Specifies the full matrix format when importing a user-specified proposal covariance.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

When importing the MCMC proposal covariance from a user-specified filename, this keyword declares the use of a full matrix format, i.e., the user provides all values of the matrix, not just the diagonal. The matrix must be symmetric, positive-definite.
Examples

method, 
bayes_calibration queso
samples = 1000 seed = 348
dram
proposal_covariance
matrix filename ‘dakota_cantilever_queso.matrix.dat’

7.2.68.6 experimental_design

• Keywords Area
• method
• bayes_calibration
• experimental_design

(Experimental) Adaptively select experimental designs for iterative Bayesian updating

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>initial_samples</td>
<td>Number of data points used during initial Bayesian calibration</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>numCandidates</td>
<td>Number of candidate design points considered</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_hifi_evaluations</td>
<td>Maximum number of high-fidelity model runs to be used</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>batch_size</td>
<td>Number of optimal designs selected concurrently</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_candidate_points_file</td>
<td>Specify text file containing candidate design points</td>
</tr>
</tbody>
</table>
Description

This "experimental design" algorithm uses responses produced by a high-fidelity model as data to update the parameters of a low-fidelity model using Bayes’ Rule. It is a capability that is under active development and is currently compatible only with queso. The user-specified high-fidelity model should depend only on configuration variables (i.e. design conditions), such as temperature or spatial location, while the user-specified low-fidelity model should depend on both configuration variables and its own model parameters to be calibrated.

The algorithm starts with a preliminary Bayesian calibration using the number of data points specified in initial_samples. These data points can be read in through the calibration_data_file command in the responses block. If num_experiments is less than initial_samples or if no such data file is provided, Latin Hypercube Samples of the design space (specified in the variables block) will be run through the user-specified high-fidelity code to supplement the initial data. Once this first calibration is complete, a set of possible experimental design conditions, specified using the configuration variables, is proposed. The user specifies the size of this set using num_candidates. The set of candidates itself may be explicitly given through the import_candidate_points_file command. If the number of candidates in this file is less than num_candidates, or if this file is omitted, the set of candidate designs will again be supplemented with a Latin Hypercube Sample of the design space.

For each candidate design $\xi_i$ in the set of possible design conditions, the mutual information (MI) between the low-fidelity model parameters $\theta$ and the high-fidelity model response $y(\xi_i)$,

$$MI = \int \int f(\theta, y(\xi)) \log \frac{f(\theta, y(\xi))}{f(\theta) f(y(\xi))} d\theta dy,$$

is approximated. The high-fidelity model is replaced by the low-fidelity model and a $k$-nearest neighbor approximation is used in the calculation. The design point $\xi^*$ for which MI is the largest is selected and run through the high-fidelity model to yield a new observation $y(\xi^*)$. This new observation is added to the calibration data, and a subsequent Bayesian calibration is performed. A new MI for each remaining candidate design is computed, and the process repeats until one of three stopping criteria are met. Multiple optimal designs may be selected concurrently by specifying batch_size.

Of the three stopping criteria, two are automatically checked by Dakota. If the relative change in the MI from one iteration to the next is sufficiently small or if the set of candidate design conditions has been exhausted, the algorithm terminates. The user may specify the third stopping criteria using max_hifi_evaluations. This limits the number of high-fidelity model evaluations that will be performed during this algorithm. It therefore limits the number of iterations through the algorithm that will be performed. Any high-fidelity model runs needed to produce the data set for the initial calibration are not included in this allocation.

In the case that the high-fidelity model must be run independently of Dakota, the user may set max_hifi_evaluations to zero. The optimal experimental design point will be calculated and reported, but the high-fidelity model will not be run. For more details, see the User’s Manual.

Expected Output

Information regarding the progress and termination condition of the experimental design algorithm is output to the screen with varying levels of verbosity. Further details can be found, regardless of verbosity, in the output file experimental_design_output.txt

Usage Tips

Due to the optional file read-ins and the supplemental sampling, it is important for the user to check consistency within the input file specifications. For example, if num_experiments is less than the number of experiments in the calibration_data_file, only the first lines of the file will be used and the rest will be discarded. The same holds true for the import_candidate_points_file and num_candidates.
Examples

The input file below illustrates the use of `experimental_design` and its options

```bash
bayes_calibration
  queso
dram
  chain_samples = 1000 seed = 348
experimental_design
  initial_samples = 5
  num_candidates = 10
  import_candidate_points_file = "bayes_experimental_design.cand.dat"
  freeform
  max_hifi_evaluations = 3
```

7.2.68.6.1 initial_samples

- **Keywords Area**
- method
- bayes_calibration
- experimental_design
- initial_samples

Number of data points used during initial Bayesian calibration

**Specification**

**Alias:** samples

**Argument(s):** INTEGER

**Default:** method-dependent

**Description**

The experimental design algorithm starts with a preliminary Bayesian calibration. The keyword `initial_samples` specifies the number of data points used during this initial calibration. These data points may come from external data sources through `calibration_data_file`, from an LHS sample of the design space, or both.

7.2.68.6.2 num_candidates

- **Keywords Area**
- method
- bayes_calibration
- experimental_design
- num_candidates

Number of candidate design points considered
Specification

**Alias**: none  
**Argument(s)**: INTEGER  
**Default**: 1

**Description**

After the initial Bayesian calibration, a set of candidate design points is created, from which the optimal design(s) will be chosen. This set is of size `num_candidates`. These candidate points may be specified by the user through `import_candidate_points_file`, from an LHS sample of the design space, or both.

7.2.68.6.3  max_hifi_evaluations

*Keywords Area*

* method

* bayes_calibration

* experimental_design

* max_hifi_evaluations

Maximum number of high-fidelity model runs to be used

**Specification**

**Alias**: none  
**Argument(s)**: INTEGER

**Description**

There are three stopping criteria by which the Bayesian experimental design algorithm will be terminated:

- The high-fidelity model has been run `max_hifi_evaluations` times
- The set of candidate design points have been exhausted
- The relative change in the mutual information from one iteration to the next is sufficiently small. The first criterion is controlled by this keyword specification

**Default Behavior**

If no maximum number of high-fidelity model runs is specified, only the last two stopping criteria listed above will be evaluated.

**Usage Tips**

Be wary that `max_hifi_evaluations` does not include any high-fidelity model evaluations that need to be run to produce the data set for the initial Bayesian calibration. This number only includes those evaluations performed on the sequentially chosen optimal design points.

If `max_hifi_evaluations` is set to zero, the a single optimal experimental design point will be selected, and Dakota will exit without running the high-fidelity model.
7.2.68.6.4  batch_size

- Keywords Area
- method
- bayes_calibration
- experimental_design
- batch_size

Number of optimal designs selected concurrently

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

*Each iteration of the experimental_design algorithm yields batch_size optimal designs.* If `max_hifi` is set to zero, these values will merely be reported in Dakota output and the file `experimental_design_output.txt`. Otherwise, the high-fidelity model will be run at all optimal designs, and the responses will be added to the list of experiment data before the Bayesian calibration is performed in the next iteration of the algorithm.

**Default Behavior**

By default, optimal designs will be selected one at a time.

7.2.68.6.5  import_candidate_points_file

- Keywords Area
- method
- bayes_calibration
- experimental_design
- import_candidate_points_file

Specify text file containing candidate design points

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**
### Description

Enables text file import of user-specified design points (i.e. configuration variables). No model responses should be included. Each row should correspond to a single design or configuration.

### Default Behavior

Be default, methods do not import points from a file.

### Usage Tips

Dakota parses input files without regard to whitespace, but the import_build_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

#### 7.2.68.6.5.1 custom_annotated

- Keywords Area
- method
- bayes_calibration
- experimental_design
- import_candidate_points_file
- custom_annotated

Selects custom-annotated tabular file format

### Topics

This keyword is related to the topics:

- file_formats

### Specification

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id   x1   x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1   0.9   1.1   0.0002   0.26   0.76
2   0.90009   1.1  0.0001996404857   0.2601620081   0.759955
3   0.89991   1.1  0.0002003604863   0.2598380081   0.760045
...```
7.2.68.6.5.2 header

• Keywords Area
• method
• bayes_calibration
• experimental_design
• import_candidate_points_file
• custom_annotated
• header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.6.5.3 eval_id

• Keywords Area
• method
• bayes_calibration
• experimental_design
• import_candidate_points_file
• custom_annotated
• eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated
7.2.68.5.4  interface_id
  
  - Keywords Area
  - method
  - bayes_calibration
  - experimental_design
  - import_candidate_points_file
  - custom_annotated
  - interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.5.5  annotated
  
  - Keywords Area
  - method
  - bayes_calibration
  - experimental_design
  - import_candidate_points_file
  - annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

  - file_formats

Specification

Alias: none
Argument(s): none
Default: annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

evironment
tabular_data
tabular_data_file = ’dakota_summary.dat’
annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...

7.2.68.6.5.6 freeform

• Keywords Area

• method

• bayes_calibration

• experimental_design

• import_candidate_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats
Specification

Alias: none  
Argument(s): none

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
  0.9  1.1  0.0002  0.26  0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

Use second Kraskov algorithm to compute mutual information

7.2.68.6.6  ksg2

- Keywords Area
  - method
  - bayes_calibration
  - experimental_design
  - ksg2
This algorithm is derived in [58]. The mutual information between $m$ random variables is approximated by

$$I_2(X_1, X_2, \ldots, X_m) = \psi(k) + (m-1)\psi(N) - (m-1)/k - <\psi(n_{x_1}) + \psi(n_{x_2}) + \ldots + \psi(n_{x_m})>,$$

where $\psi$ is the digamma function, $k$ is the number of nearest neighbors being used, and $N$ is the number of samples available for the joint distribution of the random variables. For each point $z_i = (x_{1,i}, x_{2,i}, \ldots, x_{m,i})$ in the joint distribution, $z_i$ and its $k$ nearest neighbors are projected into each marginal subspace. For each subspace $j = 1, \ldots, m$, $\varepsilon_{j,i}$ is defined as the radius of the $l_\infty$-ball containing all $k+1$ points. Then, $n_{x_{j,i}}$ is the number of points in the $j$-th subspace within a distance of $\varepsilon_{j,i}$ from the point $x_{j,i}$. The angular brackets denote that the average of $\psi(n_{x_{j,i}})$ is taken over all points $i = 1, \ldots, N$.

Examples

```
method bayes_calibration queso
dram
seed = 34785
chain_samples = 1000
posterior_stats mutual_info
ksg2
```

```
method bayes_calibration
queso
dram
chain_samples = 1000 seed = 348
experimental_design
initial_samples = 5
num_candidates = 10
max_hifi_evaluations = 3
ksg2
```

7.2.68.7 calibrate_error_multipliers

- Keywords Area
- method
- bayes_calibration
- calibrate_error_multipliers

Calibrate hyper-parameter multipliers on the observation error covariance

Specification

Alias: none
Argument(s): none
Default: none
Child Keywords:
### Description

Calibrate one or more multipliers on the user-provided observation error covariance (\texttt{experiment_variance_type}). Options include one multiplier on the whole block-diagonal covariance structure, one multiplier per experiment covariance block, one multiplier per response covariance block, or separate multipliers for each response/experiment pair (for a total of number experiments X number response groups).

**Default Behavior:** No hyper-parameter calibration. When hyper-parameter calibration is enabled, the default prior on the multiplier is a diffuse inverse gamma, with mean and mode approximately 1.0.

**Expected Output:** Final calibration results will include both inference parameters and one or more calibrated hyper-parameters.

**Usage Tips:** The per_response option can be useful when each response has its own measurement error process, but all experiments were gathered with the same equipment and conditions. The per_experiment option might be used when working with data from multiple independent laboratories.

### Examples

Perform Bayesian calibration with 2 calibration variables and two hyper-parameter multipliers, one per each of two responses. The multipliers are assumed the same across the 10 experiments. The priors on the multipliers are specified using the \texttt{hyperprior_alphas} and \texttt{hyperprior_betas} keywords.

```bash
bayes_calibration queso
  samples = 1000 seed = 348
dram
  calibrate_error_multipliers per_response
  hyperprior_alphas = 27.0
```
hyperprior_betas = 26.0

variables
uniform_uncertain 2
upper_bounds 1.e8 10.0
lower_bounds 1.e6 0.1
initial_point 2.85e7 2.5
descriptors 'E' 'w'

responses
calibration_terms = 2
calibration_data_file = 'expdata.withsigma.dat'
freeform
num_experiments = 10
experiment_variance_type = 'scalar'

7.2.68.7.1 one

- Keywords Area
- method
- bayes_calibration
- calibrate_error_multipliers
- one

Calibrate one hyper-parameter multiplier across all responses/experiments

Specification

Alias: none
Argument(s): none

Description

A single hyper-parameter multiplying all response/experiment covariance blocks will be estimated.

7.2.68.7.2 per_experiment

- Keywords Area
- method
- bayes_calibration
- calibrate_error_multipliers
- per_experiment

Calibrate one hyper-parameter multiplier per experiment

Specification

Alias: none
Argument(s): none
Description

One hyper-parameter multiplying each experiment covariance block will be estimated.

7.2.68.7.3 per_response

- Keywords Area
- method
- bayes_calibration
- calibrate_error_multipliers
- per_response

Calibrate one hyper-parameter multiplier per response

Specification

Alias: none
Argument(s): none

Description

One hyper-parameter multiplying each response covariance block will be estimated.

7.2.68.7.4 both

- Keywords Area
- method
- bayes_calibration
- calibrate_error_multipliers
- both

Calibrate one hyper-parameter multiplier for each response/experiment pair

Specification

Alias: none
Argument(s): none

Description

One hyper-parameter multiplying each experiment/response covariance block will be estimated.

7.2.68.7.5 hyperprior_alphas

- Keywords Area
- method
- bayes_calibration
• calibrate_error_multipliers
• hyperprior_alphas

Shape (alpha) of the inverse gamma hyper-parameter prior

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>hyperprior_betas</td>
<td>Scale (beta) of the inverse gamma hyper-parameter prior</td>
</tr>
</tbody>
</table>

**Description**

Shape of the prior distribution for calibrated error multipliers. Either a single value or number of hyper-parameters values may be specified.

**Default:** 102.0 (with beta = 103.0) so mean and mode are approximately 1.0 and standard deviation is about 0.1.

7.2.68.7.5.1 hyperprior_betas

• Keywords Area
• method
• bayes_calibration
• calibrate_error_multipliers
• hyperprior_alphas
• hyperprior_betas

Scale (beta) of the inverse gamma hyper-parameter prior

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Description**

Scale of the prior distribution for calibrated error multipliers. Either a single value or number of hyper-parameters values may be specified.

Defaults to 103.0 (with alpha = 102.0) so mean and mode are approximately 1.0 and standard deviation is about 0.1.
7.2.68.8 burn_in_samples

- Keywords Area
- method
- bayes_calibration
- burn_in_samples

Manually specify the burn in period for the MCMC chain.

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 0

Description

The burn in period is the number of samples at the beginning of the MCMC chain that are discarded. The resulting chain is less dependent on the starting point of the chain.

**Default Behavior**

If not specified, no MCMC samples are discarded

**Expected Output**

If `burn_in_samples` is specified, an additional tabular output file containing the final MCMC chain is generated.

Examples

Generate a chain of length 1000 and discard the first 100 points, resulting in a chain of length 900.

```plaintext
method, bayes_calibration queso
    chain_samples = 1000 seed = 348
dram
    proposal_covariance
diagonal values 1.0e6 1.0e-1
burn_in_samples 100
```

7.2.68.9 posterior_stats

- Keywords Area
- method
- bayes_calibration
- posterior_stats

Compute information-theoretic metrics on posterior parameter distribution
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Information theory allows for the calculation and quantification of the information contained within a distribution. It is particularly useful for distributions which are not Gaussian, such as those which are bimodal or highly skewed. The `posterior_stats` command calculates information metrics relating to the posterior distribution of the model parameters. These metrics are approximated by making use of the MCMC chain produced during the Bayesian update. This capability can be used with any Bayesian method.

#### 7.2.68.9.1 kl_divergence

- **Keywords Area**
  - method
  - bayes_calibration
  - posterior_stats
  - kl_divergence

Calculate the Kullback-Leibler Divergence between prior and posterior

#### Specification

- **Alias**: none
- **Argument(s)**: none

#### Description

The Kullback-Leibler (KL) Divergence, also called the relative entropy, provides a measure of the difference between two probability distributions. By specifying `kl_divergence`, the KL Divergence between the posterior \( f(\theta|y_{Data}) \) and the prior \( f(\theta) \) parameter distributions is calculated such that

\[
D_{KL} = \int f(\theta|y_{Data}) \log \frac{f(\theta|y_{Data})}{f(\theta)} d\theta
\]
This quantity can be interpreted as the amount of information gained about the parameters during the Bayesian update.

**Expected Output**

If `kl_divergence` is specified, the calculated value will be reported to the screen at the end of the calibration, following the sample statistics of the response functions. Example output is given below.

**Additional Discussion**

The quantity calculated is a $k$-nearest neighbor approximation of the possibly multi-dimensional integral given above. Therefore, some applications whose true KL Divergence is quite close to zero may report a negative KL Divergence.

**Examples**

Below is a method block of a Dakota input file that indicates the calculation of the KL Divergence

```plaintext
method,
    bayes_calibration queso
    dram
    seed = 34785
    chain_samples = 1000
    posterior_stats kl_divergence

The calculated KL approximation is indicated in the screen output by "Information gained from prior to posterior" as shown below

Sample moment statistics for each response function:

<table>
<thead>
<tr>
<th>Function</th>
<th>Mean</th>
<th>Std Dev</th>
<th>Skewness</th>
<th>Kurtosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>least_sq_term_1</td>
<td>3.9982462078e-01</td>
<td>4.7683816550e-04</td>
<td>-2.3448518080e+00</td>
<td>7.7381497770e+00</td>
</tr>
</tbody>
</table>

Information gained from prior to posterior = 1.0066819600e+01

<<<<< Iterator bayes_calibration completed.
<<<<< Environment execution completed.
```

7.2.68.9.2 **mutual_info**

- **Keywords Area**
- **method**
- **bayes_calibration**
- **posterior_stats**
- **mutual_info**

Calculate the mutual information between prior and posterior

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
## Description

The mutual information quantifies how much information two random variables contain about each other. It is a measure of the mutual dependence of two random variables. The mutual information is a non-negative measure, with zero representing complete independence of the two random variables. For continuous random variables $X$ and $Y$, the mutual information is

$$I(X,Y) = \int \int p(x,y) \log \frac{p(x,y)}{p(x)p(y)} \, dx \, dy.$$ 

The mutual information can also be interpreted as the reduction in uncertainty of one random variable due to the knowledge of another. By specifying `mutual_info`, the mutual information between the posterior parameters and the prior parameters is calculated.

The mutual information is calculated using a $k$-nearest neighbor approximation algorithm. As of Dakota 6.6, there are two such algorithms available, both of which are derived in [58]. By default, Dakota uses the first such algorithm; the second may be selected by specifying the keyword `ksg2`. Further details can be found in the Dakota Theory Manual [16].

### Expected Output

If `mutual_information` is specified, the calculated value will be reported to the screen at the end of the calibration.

### Additional Discussion

Due to the necessary approximation of the multidimensional integral above, a negative mutual information may be reported for applications whose true value is close to or equal to zero. As of Dakota 6.6, mutual information calculations are primarily used in the implementation of the `experimental_design` algorithm.

### Examples

```plaintext
method bayes_calibration queso
dram
seed = 34785
chain_samples = 1000
posterior_stats mutual_info
```

<table>
<thead>
<tr>
<th>Optional</th>
<th><code>ksg2</code></th>
<th>Use second Kraskov algorithm to compute mutual information</th>
</tr>
</thead>
</table>

- **Keywords Area**
- **method**
- **bayes_calibration**
- **posterior_stats**
- **mutual_info**
- **ksg2**
Specification

Alias: none

Argument(s): none

Description

This algorithm is derived in [58]. The mutual information between \( m \) random variables is approximated by

\[
I_2(X_1, X_2, \ldots, X_m) = \psi(k) + (m - 1)\psi(N) - (m - 1)/k - < \psi(n_{x_1}) + \psi(n_{x_2}) + \ldots + \psi(n_{x_m}) > ,
\]

where \( \psi \) is the digamma function, \( k \) is the number of nearest neighbors being used, and \( N \) is the number of samples available for the joint distribution of the random variables. For each point \( z_i = (x_{1,i}, x_{2,i}, \ldots, x_{m,i}) \) in the joint distribution, \( z_i \) and its \( k \) nearest neighbors are projected into each marginal subspace. For each subspace \( j = 1, \ldots, m, \varepsilon_{j,i} \) is defined as the radius of the \( l_\infty \)-ball containing all \( k + 1 \) points. Then, \( n_{x_{j,i}} \) is the number of points in the \( j \)-th subspace within a distance of \( \varepsilon_{j,i} \) from the point \( x_{j,i} \). The angular brackets denote that the average of \( \psi(n_{x_{j,i}}) \) is taken over all points \( i = 1, \ldots, N \).

Examples

```dakota
method
  bayes_calibration queso
dram
  seed = 34785
  chain_samples = 1000
  posterior_stats mutual_info
  ksg2

method
  bayes_calibration
  queso
dram
  chain_samples = 1000 seed = 348
  experimental_design
  initial_samples = 5
  num_candidates = 10
  max_hifi_evaluations = 3
  ksg2
```

```
7.2.68.9.3 kde
  • Keywords Area
  • method
  • bayes_calibration
  • posterior_stats
  • kde

Calculate the Kernel Density Estimate of the posterior distribution
```

Specification

Alias: none

Argument(s): none
Description

A kernel density estimate (KDE) is a non-parametric, smooth approximation of the probability density function of a random variable. It is calculated using a set of samples of the random variable. If $X$ is a univariate random variable with unknown density $f$ and independent and identically distributed samples $x_1, x_2, \ldots, x_n$, the KDE is given by

$$
\hat{f} = \frac{1}{nh} \sum_{i=1}^{n} K\left(\frac{x - x_i}{h}\right).
$$

The kernel $K$ is a non-negative function which integrates to one. Although the kernel can take many forms, such as uniform or triangular, Dakota uses a normal kernel. The bandwidth $h$ is a smoothing parameter that should be optimized. Choosing a large value of $h$ yields a wide KDE with large variance, while choosing a small value of $h$ yields a choppy KDE with large bias. Dakota approximates the bandwidth using Silverman’s rule of thumb,

$$
h = \hat{\sigma} \left(\frac{4}{3n}\right)^{1/5},
$$

where $\hat{\sigma}$ is the standard deviation of the sample set $\{x_i\}$.

For multivariate cases, the random variables are treated as independent, and a separate KDE is calculated for each.

Expected Output

If kde is specified, calculated values of $\hat{f}$ will be output to the file kde_posterior.dat. Example output is given below.

Examples

Below is a method block of a Dakota input file that indicates the calculation of the KDE

```dakota
method,
  bayes_calibration queso
    dram
    seed = 34785
    chain_samples = 1000
    posterior_stats kde
```

The calculated KDE values are output to the file kde_posterior.dat, as shown below

```
uuv_1  KDE PDF estimate
  0.406479 61.2326
  0.406338 64.0245
  0.402613 114.468
  0.402613 114.468
  0.40249 114.409
  0.40282 114.162
  0.398899 65.2361
  0.400093 84.9285
  0.401264 104.105
  0.400917 98.7803
```

7.2.68.10 chain_diagnostics

- Keywords Area
- method
- bayes_calibration
- chain_diagnostics

Compute diagnostic metrics for Markov chain
Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>confidence_intervals</td>
<td>Calculate the confidence intervals on estimates of first and second moments</td>
</tr>
</tbody>
</table>

Description

While a Markov chain produced via Monte Carlo sampling eventually converges to a set of samples representative of an underlying probability distribution, the first set of samples may not accurately capture the target distribution. Chain diagnostic metrics provide measures of this convergence, and are intended to help users decide whether samples should be increased in Bayesian calibration exercises.

7.2.68.10.1 confidence_intervals

- Keywords Area
- method
- bayes_calibration
- chain_diagnostics
- confidence_intervals

Calculate the confidence intervals on estimates of first and second moments

Specification

Alias: none
Argument(s): none

Description

During Bayesian calibration, a chain of samples is produced, which represents the underlying posterior distribution of model parameters. For each parameter sample, the corresponding model response is computed. The confidence_intervals keyword indicates the calculation of a 95% confidence interval on the estimated mean and variance of each parameter and each response.

As of Dakota 6.10, these confidence intervals are calculated using the asymptotically valid interval estimator,

$$\bar{g}_n \pm t_\alpha \frac{\hat{\sigma}_n}{\sqrt{n}}$$

where $\bar{g}_n$ is the moment (i.e. mean or variance), $t_\alpha$ is the Student’s $t$-value for the 95th quantile, $n$ is the sample size, and $\hat{\sigma}_n$ is an estimate of the standard error whose square is obtained using batch means estimation. The Markov chain produced via Monte Carlo sampling eventually converges to a set of samples representative of an underlying probability distribution, the first set of samples may not accurately capture the target distribution. Chain diagnostic metrics provide measures of this convergence, and are intended to help users decide whether samples should be increased in Bayesian calibration exercises.

Markov chain produced via Monte Carlo sampling eventually converges to a set of samples representative of an underlying probability distribution, the first set of samples may not accurately capture the target distribution. Chain diagnostic metrics provide measures of this convergence, and are intended to help users decide whether samples should be increased in Bayesian calibration exercises.

As of Dakota 6.10, these confidence intervals are calculated using the asymptotically valid interval estimator,$$\bar{g}_n \pm t_\alpha \frac{\hat{\sigma}_n}{\sqrt{n}}$$where $\bar{g}_n$ is the moment (i.e. mean or variance), $t_\alpha$ is the Student’s $t$-value for the 95th quantile, $n$ is the sample size, and $\hat{\sigma}_n$ is an estimate of the standard error whose square is obtained using batch means estimation. The Markov chain produced via Monte Carlo sampling eventually converges to a set of samples representative of an underlying probability distribution, the first set of samples may not accurately capture the target distribution. Chain diagnostic metrics provide measures of this convergence, and are intended to help users decide whether samples should be increased in Bayesian calibration exercises.

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chain produced during calibration is broken up into “batches,” the sample moment is calculated for each batch, and \( \hat{\sigma}_n \) is an unbiased estimate of the standard deviation in these batch moment calculations.

**Expected Output**

If `confidence_intervals` is specified, the 95% confidence interval for the mean and variance for each parameter and for each response will be output to the screen. If `output` is set to `debug`, the mean of the moment calculated for each batch will also be output to the screen.

**Additional Discussion**

Confidence intervals may be used to indicate to the user whether samples needs to be increased during the Bayesian calibration. For example, if the width of the intervals (one, many, or all) is below some threshold value, that may indicate that enough samples have been drawn.

**Examples**

Below is a method block of a Dakota input file that indicates the calculation of confidence intervals

```
method,
    bayes_calibration queso
dram
    seed = 34785
chain_samples = 1000
chain_diagnostics
    confidence_intervals
```

The calculated confidence intervals are output to the screen under “Chain diagnostics”:

```
Sample moment statistics for each posterior variable:
    Mean  Std Dev  Skewness  Kurtosis
e  2.8609959149e+07  1.4417714265e+05  8.0289072766e-01  7.8655956160e-02
w  2.5016445558e+00  3.8306697138e-03  -1.2217188066e-01  3.8866929786e-02

Sample moment statistics for each response function:
    Mean  Std Dev  Skewness  Kurtosis
stress  2.6282814617e+03  8.9765361327e+01  1.3400226598e+01  4.9239052296e-02
displacement  2.9604502307e-01  1.0636886950e-02  -3.5080744509e-01  -1.2381836901e-01

Chain diagnostics
    95% Confidence Intervals of means
    E = [2.8570364780e+07, 2.8649553519e+07]
    w = [2.5009524557e+00, 2.5023366559e+00]
    stress = [2.6120609285e+03, 2.6445019948e+03]
    displacement = [2.9337418438e-01, 2.9871586175e-01]
    95% Confidence Intervals of variances
    E = [1.5074828429e+10, 2.6499268497e+10]
    w = [1.1359880885e-05, 1.7988180028e-05]
    stress = [6.2340446164e+03, 9.8815955721e+03]
    displacement = [8.8187472572e-05, 1.3809925539e-04]
```

**7.2.68.11 model_evidence**

- **Keywords Area**
  - method
  - bayes_calibration
  - model_evidence

Calculate model evidence (marginal likelihood of model) when using Bayesian methods
Model evidence is used in Bayesian model selection and model averaging. It is defined as the probability of the data given the model, and is calculated by averaging the likelihood of the model parameters over all values of the model parameters according to their prior distributions. In Dakota, one must calculate the model evidence separately for each model and perform the normalization to obtain the posterior model plausibility for each model.

**Default Behavior**

When specifying `model_evidence`, there are two methods of calculating it. One or both may be specified. They include the Monte Carlo approximation, given by `mc_approx` and the Laplace approximation, given by `laplace_approx`. `mc_approx` is the default approach.

**Expected Output** Currently, the model evidence will be printed in the screen output with prefacing text indicating if it is calculated by Monte Carlo sampling or the Laplace approximation.

**Usage Tips**

- `7.2.68.11.1 mc_approx`
  - Keywords Area
  - method
  - bayes_calibration
  - model_evidence
  - mc_approx

Calculate model evidence using a Monte Carlo sampling approach
The `mc_approx` keyword for model evidence indicates that sample values will be generated from the prior distribution, and then the simulation model will be evaluated at these sample values to obtain corresponding likelihood values. The average of the likelihood weighted by the prior is the model evidence. The accuracy of this approximation depends on the number of samples taken, which is specified by the `evidence_samples` keyword.

**Default Behavior**

If `evidence_samples` is not specified with `mc_approx`, Dakota uses the number of chain samples from the MCMC (`chain_samples`) as the number of samples to use for calculating the model evidence.

**Expected Output**

Currently, the model evidence will be printed in the screen output with prefacing text indicating if it is calculated by Monte Carlo sampling.

**Usage Tips**

- **Keywords Area**
  - method
  - bayes_calibration
  - model_evidence
  - evidence_samples

The number of samples used in Monte Carlo approximation of the model evidence.

**Specification**

- **Alias**: none
- **Argument(s)**: INTEGER

**Description**

The `mc_approx` keyword for model evidence indicates that sample values will be generated from the prior distribution, and then the simulation model will be evaluated at these sample values to obtain corresponding likelihood values. The average of the likelihood weighted by the prior is the model evidence. The accuracy of this approximation depends on the number of samples taken, which is specified by the `evidence_samples` keyword. Note that each sample specified by the `evidence_samples` keyword will require an evaluation of the simulation model to compute the corresponding likelihood. So, this may become costly for expensive simulations. Additionally, many prior samples will have very low (near zero) likelihood, so millions of samples may be required for accurate computation of the integral which defines model evidence.

**Default Behavior**

If `evidence_samples` is not specified with `mc_approx`, Dakota uses the number of chain samples from the MCMC (`chain_samples`) as the number of samples to use for calculating the model evidence.

**Expected Output**

Currently, the model evidence will be printed in the screen output with prefacing text indicating if it is calculated by Monte Carlo sampling.

**Usage Tips**
7.2.68.11.3 laplace_approx

- Keywords Area
- method
- bayes_calibration
- model_evidence
- laplace_approx

Calculate model evidence using the Laplace approximation

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `laplace_approx` keyword for model evidence indicates that a pre-solve will be used prior to the Bayesian MCMC sampling to estimate the Maximum A Posteriori (MAP) point. The Laplace approximation assumes the posterior density is nearly Gaussian and is given by a formula which involves the likelihood at the MAP point, the prior density at the MAP point, and the Hessian of the log-posterior at the MAP point. The formula is given in the Dakota User's manual. This method is efficient at estimating the model evidence for posterior densities with weak non-Gaussian characteristics but it does require a MAP solve (so `pre-solve` should be specified) and it does require gradient and Hessians of the response to be on.

**Default Behavior**

**Expected Output**  Currently, the model evidence will be printed in the screen output with prefacing text indicating if it is calculated by the Laplace approximation.

**Usage Tips**

7.2.68.12 model_discrepancy

- Keywords Area
- method
- bayes_calibration
- model_discrepancy

(Experimental) Post-calibration calculation of model discrepancy correction

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
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<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
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<td>discrepancy_type</td>
<td></td>
<td>Specify the type of model discrepancy</td>
</tr>
<tr>
<td>Optional</td>
<td>num_prediction_configs</td>
<td></td>
<td>Specify number of prediction locations for model discrepancy</td>
</tr>
<tr>
<td>Optional</td>
<td>prediction_configs</td>
<td></td>
<td>List prediction locations for model discrepancy</td>
</tr>
<tr>
<td>Optional</td>
<td>import_prediction_configs</td>
<td></td>
<td>Specify text file containing prediction configurations for model discrepancy</td>
</tr>
<tr>
<td>Optional</td>
<td>export_discrepancy_file</td>
<td></td>
<td>Output file for prediction discrepancy calculations</td>
</tr>
<tr>
<td>Optional</td>
<td>export_corrected_model_file</td>
<td></td>
<td>Output file for corrected model prediction calculations</td>
</tr>
<tr>
<td>Optional</td>
<td>export_corrected_variance_file</td>
<td></td>
<td>Output file for prediction variance calculations</td>
</tr>
</tbody>
</table>

**Description**

The goal of parameter calibration is to minimize the difference between experimental data, \(d(x)\), and model observations, \(M(\theta, x)\), where \(\theta\) are the model parameters and \(x\) is a configuration variable, such as temperature or pressure. However, it is not uncommon that, at the conclusion of parameter calibration, the agreement between experimental data and the calibrated model is not "close enough." This is often due to model form or structural error. In this case, the canonical equation

\[
d(x) = M(\theta, x) + \varepsilon
\]

is replaced by one that also includes model discrepancy \(\delta(x)\),

\[
d(x) = M(\theta, x) + \delta(x) + \varepsilon.
\]

In the Dakota implementation, the calculation of \(\delta(x)\) is performed after the model parameters \(\theta\) are calibrated. For each observable \(d_i\), the discrepancy

\[
\delta_i(x_j) = d_i(x_j) - M_i(\theta^*, x_j)
\]

is calculated for each value \(x_j\) of the configuration variable, where \(\theta^*\) is the optimal parameter value obtained during the calibration. For scalar responses, the model discrepancy is only a function of the configuration variables, and
there is one discrepancy regression model for each observable $d_i$. This set of discrepancy models may be specified to be either Gaussian process or polynomial regression models of constant, linear, or quadratic order, and each model is fit to the calculated discrepancy values. See the `discrepancy_type` command for more details regarding these options. For field responses, the model discrepancy is a function of the configuration variables as well as the independent field coordinates (such as time or space), and there is one discrepancy regression model for each field. In this case, the discrepancy models are Gaussian process models. The calculation of model discrepancy has not been tested for cases in which responses are mixed scalar and field responses.

The user may then specify new or "prediction" configurations at which the corrected model $M(\theta^*, x_{new}) + \delta(x_{new})$ should be calculated, using one of the `num_prediction_configs`, `prediction_configs`, or `import_prediction_configs` keywords. If none of these keywords is specified, the number of prediction configurations is set to 10 for scalar responses. The corresponding prediction variances are also calculated, according to

$$\Sigma_{total}(x) = \Sigma_{\delta}(x) + \sigma^2_{\text{exp}}.$$  

Here, $\Sigma_{\delta}(x)$ is the (co)variance calculated from the Gaussian process or polynomial regression model, and $\sigma^2_{\text{exp}}$ is the maximum variance, taken over all configuration variables for that observation. In the case of field responses, the default prediction configurations are set equal to the input configurations, and the variance information contains only the variance calculated from the Gaussian process correction model. Further details can be found in the Dakota User's and Theory Manuals.

**Usage Tips**

For field responses, the keyword `read_field_coordinates` must be specified when computing the model discrepancy. See `field_calibration_terms` for more information regarding options for calibration with field responses.

**Expected Output**

The resulting values of $\delta(x_{new})$ will be exported to the file specified using `export_discrepancy_file` or to the default file `dakota_discrepancy_tabular.dat`. The values of the corrected model at the specified prediction configurations will be exported to the file specified using `export_corrected_model_file` or to the default file `dakota_corrected_model_tabular.dat`, and the corresponding prediction variances will be exported to `dakota_discrepancy_variance_tabular.dat` or the file specified with `export_corrected_variance_file`.

**Examples**

Extensive examples can be found in the Dakota User's and Theory Manuals. The input files below illustrate the options available to `model_discrepancy`

```plaintext
model_discrepancy
  discrepancy_type  gaussian_process
  num_prediction_configs  11
  export_discrepancy_file  "discrepancy_values.txt"
  export_corrected_model_file  "corrected_model.txt"
  export_corrected_variance_file  "prediction_variance.txt"

model_discrepancy
  discrepancy_type  polynomial
  correction_order  constant
  prediction_configs  1  1.5  2  2.5  3

model_discrepancy
  discrepancy_type  polynomial
  correction_order  linear
  import_prediction_configs  "prediction_configs.txt"
```

7.2.68.12.1 discrepancy_type
• method
• bayes_calibration
• model_discrepancy
• discrepancy_type

Specify the type of model discrepancy

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** gaussian process

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<td><strong>gaussian_process</strong></td>
<td>Use the Surfpack version of Gaussian process as the discrepancy model</td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>polynomial</strong></td>
<td>Use a polynomial surrogate as the discrepancy model</td>
</tr>
</tbody>
</table>

**Description**

After the model parameters are calibrated, the difference between the data and the calibrated model, i.e. the model discrepancy, is calculated

\[ \delta_i(x_j) = d_i(x_j) - M_i(\theta^*, x_j). \]

Each \( \delta_i \) corresponds to a different regression model. These regression models must all be either Gaussian process or polynomial models, and they are functions of the configuration variable \( x \). The order of the trend function may be selected using the `correction_order` command by specifying `constant`, `linear`, or `quadratic`.

Note that for Dakota 6.9 and earlier, this keyword only applies to discrepancy calculations for scalar responses. For field responses, a Gaussian process model with a quadratic function is used by default.

**7.2.68.12.1.1 gaussian_process**

• Keywords Area
  • method
  • bayes_calibration
  • model_discrepancy
  • discrepancy_type
  • gaussian_process

Use the Surfpack version of Gaussian process as the discrepancy model
Specification

Alias: kriging
Argument(s): none

Child Keywords:

<table>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description of Group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>trend_order</td>
<td>Trend function order of the model discrepancy</td>
</tr>
</tbody>
</table>

Description

This keyword specifies a generic version of the Gaussian process implemented in the surface-fitting library called Surfpack. With the exception of the correction_order specification, all default options are used in the approximation of the model discrepancy function. That is, DIRECT is used as the optimization method for finding the maximum likelihood estimates of the hyper-parameters that govern the trend and correlation functions, the correlation lengths are optimized by Surfpack, and ill-conditioning is handled internally; gradient-enhanced kriging is not provided as an option in this application.

By selecting this option, each discrepancy function will take the form of a Gaussian process

7.2.68.12.1.2 trend_order

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- discrepancy_type
- gaussian_process
- trend_order

Trend function order of the model discrepancy

Specification

Alias: none
Argument(s): INTEGER
Default: 2

Description

The order of the trend function may be specified using the trend_order command, followed by either 0, for which a constant trend function will be used, 1, for which a linear trend function will be used, or 2, for which a quadratic trend function will be used.

Default Behavior
By default, a quadratic trend function is used.
7.2.68.12.1.3 polynomial

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- discrepancy_type
- polynomial

Use a polynomial surrogate as the discrepancy model

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>basis_order</td>
<td>Basis function order of the model discrepancy</td>
</tr>
</tbody>
</table>

**Description**

For the purposes of model discrepancy, constant, linear, and quadratic polynomial surrogate models are available. These models are less expensive to calculate than Gaussian process models but are consequently less flexible to discrepancy trends that are not constant, linear, or quadratic in nature. For more information, see the Dakota User’s and Theory Manuals.

By specifying this command, each discrepancy function will take the form of a polynomial regression. They will all have the same order.

7.2.68.12.1.4 basis_order

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- discrepancy_type
- polynomial
- basis_order

Basis function order of the model discrepancy
Specification

Alias: none
 Argument(s): INTEGER
 Default: 2

Description

The order of the polynomial for the model discrepancy may be specified using the `basis_order` command, followed by either 0, for which a constant polynomial will be used, 1, for which a linear polynomial will be used, or 2, for which a quadratic polynomial will be used.

Default Behavior

By default, a quadratic polynomial is used.

7.2.68.12.2 num_prediction_configs

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- num_prediction_configs

Specify number of prediction locations for model discrepancy

Specification

Alias: none
 Argument(s): INTEGER
 Default: 0

Description

Following the calculation of the model discrepancy functions, $\delta_i$, and the corrected model $M_i + \delta_i$, may be calculated at new or "prediction" configurations. Using the `num_prediction_configurations` command, the user may specify the number of prediction configurations they wish to be computed. The new configuration locations will be spread uniformly in the configuration variable domain specified in the `variables` block of the input file.

Usage Tips

If none of `num_prediction_configs`, `prediction_configs`, or `import_prediction_configs` is specified, the number of prediction configurations is set to 10 for scalar responses. For field responses, the corrected model will be output at the input values of the configuration variables.

7.2.68.12.3 prediction_configs

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
• prediction_configs

List prediction locations for model discrepancy

Specification

Alias: none
Argument(s): REALLIST

Description

Following the calculation of the model discrepancy functions, $\delta$, and the corrected model $M_i + \delta$ may be calculated at new or “prediction” configurations. Using the prediction_configs command, the user may explicitly list the locations of the prediction configurations they wish to be computed. See parent command for an example.

Usage Tips

If none of num_prediction_configs, prediction_configs, or import_prediction_configs is specified, the number of prediction configurations is set to 10 for scalar responses. For field responses, the corrected model will be output at the input values of the configuration variables.

7.2.68.12.4 import_prediction_configs

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• import_prediction_configs

Specify text file containing prediction configurations for model discrepancy

Specification

Alias: none
Argument(s): STRING
Default: no point import from a file

Child Keywords:

<table>
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<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
</tbody>
</table>
Description

Enables text file import of user-specified prediction configurations at which to calculate the corrected model $M_i + \Delta_i$.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_prediction_configs` file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.68.12.4.1 custom_annotated

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- import_prediction_configs
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1     x2     obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.12.4.2 header

- Keywords Area

- method

- bayes_calibration

- model_discrepancy
Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.12.4.3 eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.12.4.4 interface_id

Enable interface ID column in custom-annotated tabular file
7.2.68.12.4.5 annotated

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• import_prediction_configs
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none
Argument(s): none
Default: annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading `eval_id` and `interface_id` columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```
%eval_id interface   x1  x2  obj fn  nln_ineq_con_1  nln_ineq_con_2
1  NO_ID     0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```

7.2.68.12.4.6 freeform

• Keywords Area

• method

• bayes_calibration

• model_discrepancy

• import_prediction_configs

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

**Alias:** none

**Argument(s):** none

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**
The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.68.12.5 export_discrepancy_file

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_discrepancy_file

Output file for prediction discrepancy calculations

Specification

- **Alias**: none
- **Argument(s)**: STRING
- **Default**: discrepancy export to default filename
- **Child Keywords**:

Dakota Version 6.15 Reference Manual generated on November 5, 2021
### Description

Enables the user to specify the name and format of the file to which prediction discrepancy information is output. The file will contain the values of the discrepancy model $\delta_i$ calculated at those points specified with `num_predictions`, `prediction_configs`, or `import_prediction_configs`

**Default Behavior**

If `export_discrepancy_file` is not specified, information will be automatically output to `dakota_discrepancy_tabular.dat`.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**7.2.68.12.5.1 custom_annotated**

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_discrepancy_file
- custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- fileFormats

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

---

<table>
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<tr>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
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<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>
### Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1   x2   obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...```
7.2.68.12.5.2  header
  • Keywords Area
  • method
  • bayes_calibration
  • model_discrepancy
  • export_discrepancy_file
  • custom_annotated
  • header

Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

See description of parent custom_annotated

7.2.68.12.5.3  eval_id
  • Keywords Area
  • method
  • bayes_calibration
  • model_discrepancy
  • export_discrepancy_file
  • custom_annotated
  • eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

See description of parent custom_annotated
7.2.68.12.5.4  interface_id

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_discrepancy_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent custom_annotated

7.2.68.12.5.5  annotated

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_discrepancy_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated
```

Resulting tabular file:

```bash
%eval_id interface x1  x2     obj_fn  nln_ineq_con_1  nln_ineq_con_2
1 NO_ID   0.9  1.1       0.0002  0.26      0.76
2 NO_ID   0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3 NO_ID   0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...```

7.2.68.12.5.6 freeform

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_discrepancy_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
### 7.2 method

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated

### Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

### Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

### Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76
0.90009  1.1  0.0001996404857  0.2601620081  0.759955
0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

### 7.2.68.12.6 export_corrected_model_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **model_discrepancy**
- **export_corrected_model_file**

Output file for corrected model prediction calculations
Specification

Alias: none

Argument(s): STRING

Default: corrected model export to default filename

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Enables the user to specify the name and format of the file to which corrected model $M_i + \delta_i$ information is output. The file will contain the values of the discrepancy model $\delta_i$ calculated at those points specified with num_predictions, prediction_configs, or import_prediction_configs.

Corresponding variance information is also output to file, see export_corrected_variance_file

Default Behavior

If export_corrected_model_file is not specified, information will be automatically output to dakota_corrected_model_tabular.dat.

Usage Tips

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.68.12.6.1 custom_annotated

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_model_file
- custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.90</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.68.12.6.2 header

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_model_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

See description of parent custom_annotated

7.2.68.12.6.3 eval_id

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_model_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none
Description

See description of parent custom_annotated

7.2.68.12.6.4 interface_id

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• export_corrected_model_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated

7.2.68.12.6.5 annotated

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• export_corrected_model_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```
environment
  tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2.68.12.6.6 freeform

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_model_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats
### 7.2 method

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** annotated

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
  0.9   1.1   0.0002   0.26   0.76
  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
```

7.2.68.12.7 export_corrected_variance_file

- **Keywords Area**
- **method**
- **bayes_calibration**
- **model_discrepancy**
- **export_corrected_variance_file**

Output file for prediction variance calculations
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** corrected model variance export to default filename

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional(Choose One)</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

Enables the user to specify the name and format of the file to which corrected model variance information is output. The file will contain the values of the variances computed at those points specified with `num_predictions`, `prediction_configs`, or `import_prediction_configs`.

For scalar responses, the prediction variance for each observable $d_i$ is calculated according to

$$
\Sigma_{total}(x) = \Sigma_{d}(x) + \sigma^{2}_{exp}.
$$

Here, $\Sigma_{d}(x)$ is the (co)variance calculated from the Gaussian process or polynomial regression model, details of which can be found in the Dakota User’s and Theory Manuals. If experimental variance information is provided for the calibration, the maximum variance found for the observable $d_i$ is used as $\sigma^{2}_{exp}$. For more details, see the Dakota User’s and Theory Manuals. For field responses, the variance of the discrepancy model alone is output to this file.

**Default Behavior**

If `export_corrected_variance_file` is not specified, information will be automatically output to `dakota_discrepancy_variance_tabular.dat`.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.68.12.7.1 custom_annotated

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_variance_file
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1   0.9 1.1 0.0002  0.26  0.76
2   0.90009  1.1 0.0001996404857  0.2601620081  0.759955
3   0.89991  1.1 0.0002003604863  0.2598380081  0.760045
...
```

### 7.2.68.12.7.2 header

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_variance_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

See description of parent `custom_annotated`

### 7.2.68.12.7.3 eval_id

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_variance_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file
7.2 method

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.12.7.4 interface_id

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• export_corrected_variance_file
• custom_annotated
• interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.68.12.7.5 annotated

• Keywords Area
• method
• bayes_calibration
• model_discrepancy
• export_corrected_variance_file
• annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats
Specification

Alias: none
Argument(s): none
Default: annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id
- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.2.68.12.7.6 freeform

- Keywords Area
- method
- bayes_calibration
- model_discrepancy
- export_corrected_variance_file
- freeform

Selects freeform file format.
7.2 method 3081

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform

Resulting tabular file:

          0.9   1.1  0.0002   0.26    0.76
  0.90009   1.1  0.0001996404857  0.2601620081  0.759955
  0.89991   1.1  0.0002003604863  0.2598380081  0.760045
  ...
```

7.2.68.13 sub_sampling_period

- Keywords Area
Specify a sub-sampling of the MCMC chain

### Specification

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 0

### Description

If a sub_sampling_period is specified, the MCMC chain is filtered such that only the sample at the beginning of each period is in the final MCMC chain. The sub_sampling_period should therefore be greater than or equal to the correlation length of the samples.

### Default Behavior

If not specified, all MCMC samples are included in the final chain.

### Expected Output

If specified, an additional tabular output file containing the final (sub-sampled) MCMC chain is generated.

### Examples

Generate and then filter a 1000 sample MCMC chain, retaining the first accepted sample and every 100th sample thereafter.

An example method block of a Dakota input file is given below

```dakota
method,  
  bayes_calibration queso  
    chain_samples = 1000 seed = 348  
    dram  
    proposal_covariance  
      diagonal values 1.0e6 1.0e-1  
    sub_sampling_period 100

```

### 7.2.68.14 probability_levels

- **Keywords Area**
  - **method**
  - **bayes_calibration**
  - **probability_levels**

Specify probability levels at which to compute credible and prediction intervals
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_probability_levels</td>
<td>This capability is currently inactive</td>
</tr>
</tbody>
</table>

**Description**

Credible and prediction intervals of model responses are computed for specified probabilities. Credible intervals are calculated from the response function values corresponding to the final MCMC chain. Calculation of prediction intervals consider these response values as well as the experimental uncertainty, which is specified by the user via the `experiment_variance_type` command.

**Expected Output**

If `probability_levels` is specified, Dakota will create a table containing the credible intervals for each response function. The corresponding table containing the prediction intervals will also be created if a `experiment_variance_type` has been specified. This information is output to the screen and to a file. In addition, the output file contains the means and standard deviations of each response function and Gaussian approximations of the 5/95 credible and prediction intervals, in which the lower bound is two standard deviations below the mean and the upper bound is two standard deviations above the mean.

**Usage Tips**

Only one probability level needs to be specified for each desired interval. Both corresponding end points of the intervals are automatically calculated. For example, if 0.05 is specified, both the 0.05 and 0.95 probability levels are output to the screen and output file.

**Additional Discussion**

Credible intervals propagate uncertainties in parameter density information to the quantity of interest and quantify how well the model fits the provided data. Prediction intervals propagate both parameter and experimental measurement uncertainties and contain the next experimental or simulated observation with the specified probability.

**Examples**

Below is a Dakota input file specifying the calculation of credible and prediction intervals:

```dakota
method,
    bayes_calibration queso
    chain_samples = 1000 seed = 348
dram
proposal_covariance
diagonal values 1.0e6 1.0e-1
probability_levels 0.05 0.1
    0.075 0.1
variables,
    uniform_uncertain 2
    upper_bounds 1.e8 10.0
    lower_bounds 1.e6 0.1
    initial_point 2.85e7 2.5
descriptors 'E' 'w'
continuous_state 4
    initial_state 3 40000 500 1000
descriptors 't' 'R' 'X' 'Y'
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
The resulting screen output below shows the table of credible and prediction intervals.

### Credibility Intervals for stress

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.476409695e+03</td>
<td>5.000000000e-02</td>
</tr>
<tr>
<td>2.8242874802e+03</td>
<td>9.500000000e-01</td>
</tr>
<tr>
<td>2.439068791e+03</td>
<td>1.000000000e-01</td>
</tr>
<tr>
<td>2.7952985803e+03</td>
<td>9.000000000e-01</td>
</tr>
</tbody>
</table>

### Credibility Intervals for displacement

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7409870925e-01</td>
<td>7.500000000e-02</td>
</tr>
<tr>
<td>3.0891296255e-01</td>
<td>9.250000000e-01</td>
</tr>
<tr>
<td>2.7538816802e-01</td>
<td>1.000000000e-01</td>
</tr>
<tr>
<td>3.0889319332e-01</td>
<td>9.000000000e-01</td>
</tr>
</tbody>
</table>

### Prediction Intervals for stress

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.0964882850e+03</td>
<td>5.000000000e-02</td>
</tr>
<tr>
<td>3.193026765e+03</td>
<td>9.500000000e-01</td>
</tr>
<tr>
<td>2.1822183238e+03</td>
<td>1.000000000e-01</td>
</tr>
<tr>
<td>3.1099058450e+03</td>
<td>9.000000000e-01</td>
</tr>
</tbody>
</table>

### Prediction Intervals for displacement

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.355036055e-01</td>
<td>7.500000000e-02</td>
</tr>
<tr>
<td>3.509741218e-01</td>
<td>9.250000000e-01</td>
</tr>
<tr>
<td>2.4016170870e-01</td>
<td>1.000000000e-01</td>
</tr>
<tr>
<td>3.4701712866e-01</td>
<td>9.000000000e-01</td>
</tr>
</tbody>
</table>

#### 7.2.68.14.1 num_probability_levels

- **Keywords Area**
- **method**
- **bayes_calibration**
- **probability_levels**
- **num_probability_levels**

This capability is currently inactive

### Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** probability_levels evenly distributed among response functions
7.2.68.15 convergence_tolerance

- Keywords Area
- method
- bayes_calibration
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.68.16 max_iterations

- Keywords Area
- method
- bayes_calibration
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max_function_evaluations

7.2.68.17  model_pointer

- Keywords Area
- method
- bayes_calibration
- model_pointer

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- block_pointer
7.2 method

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                 0.1 0.2 0.6
                 0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
```
Turn on scaling for variables, responses, and constraints

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none

Argument(s): none

Default: no scaling

Description

Some optimization and calibration methods support scaling of continuous design variables, objective functions, calibration terms, and constraints. This is activated with the scaling keyword. Discrete variable scaling is not supported.

When scaling is enabled, variables, functions, gradients, Hessians, etc., are transformed such that the method iterates in scaled variable space, whereas evaluations of the computational model as specified in the interface are performed on the original problem scale. Therefore using scaling does not require rewriting the interface to the simulation code.

Scaling also requires the specification of additional keywords scale_types and/or *scales in the variables and responses blocks. When the scaling keyword is omitted from method, these scaling type and value specifications are ignored in the variables and responses sections.

This page describes the usage of all scaling related keywords. The additional keywords come in pairs, one pair for each set of quantities (variables, primary responses, or constraints) to be scaled.

- a *scales keyword, which gives characteristic values (divisors)
- a *scale_types keyword, which determines how to use the characteristic values

The pair of keywords both take argument(s), and the length of the arguments can be either be one or equal to the number of quantities to be scaled (see details in responses for lengths when field responses are active). If one argument is given, it will apply to all quantities in the set. See the examples below.

Scale Types

There are three scale types:
1. **value** - characteristic value scaling
   The target quantity will be scaled (divided) by the specified characteristic value.

2. **auto** - automatic scaling
   First the quantity is scaled by any characteristic value, then automatic scaling will be attempted according to the following scheme:
   - two-sided bounds scaled into the interval [0,1];
   - one-sided bound or targets are scaled by the characteristic value, moving the bound or target to 1 and changing the sense of inequalities where necessary;
   - no bounds or targets: no automatic scaling possible, therefore no scaling for this component

   Automatic scaling is not available for objective functions nor calibration terms since they lack bound constraints. Further, when automatically scaled, linear constraints are scaled by characteristic values only, not affinely scaled into [0,1].

3. **log** - logarithmic scaling
   First, any characteristic values from the optional * scales specification are applied. Then logarithm base 10 scaling is applied.
   - Logarithmic scaling is not available for linear constraints.
   - When continuous design variables are log scaled, linear constraints are not allowed.

**Scales**

The *scales keywords are used to specify the characteristic values. These must be non-zero real numbers. The numbers are used according to the corresponding *scale_types, as described above. A value of 1.0 can be used to selectively omit some quantities in a vector from being scaled.

Depending on the scale type, the characteristic values may be required or optional.

- auto, log - optional
- value - required.

A warning is issued if scaling would result in division by a value smaller in magnitude than 1.0e10*DBL_MIN. User-provided values violating this lower bound are accepted unaltered, whereas for automatically calculated scaling, the lower bound is enforced.

**Examples**

The two examples below are equivalent:

```plaintext
responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value"
  primary_scales = 1 1 100

responses
  objective_functions 3
  sense "maximize"
  primary_scale_types = "value" "value" "value"
  primary_scales = 1 1 100
```

---

**7.2.69 dace**

- Keywords Area
- method
- dace

Design and Analysis of Computer Experiments
This keyword is related to the topics:

- package_ddace
- design_and_analysis_of_computer_experiments

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td><strong>DACE type (Group 1)</strong></td>
<td>grid</td>
<td><strong>random</strong> Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>oas</td>
<td>Orthogonal Array Sampling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>lhs</td>
<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>oa_lhs</td>
<td>Orthogonal Array Latin Hypercube Sampling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>box_behnken</td>
<td>Box-Behnken Design</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central_composite</td>
<td>Central Composite Design</td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>samples</td>
<td>Number of samples for sampling-based methods</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>seed</td>
<td>Seed of the random number generator</td>
<td></td>
</tr>
<tr>
<td><strong>Optional</strong></td>
<td>fixed_seed</td>
<td>Reuses the same seed value for multiple random sampling sets</td>
<td></td>
</tr>
</tbody>
</table>
The Distributed Design and Analysis of Computer Experiments (DDACE) library provides the following DACE techniques:

1. grid sampling (grid)
2. pure random sampling (random)
3. orthogonal array sampling (oas)
4. latin hypercube sampling (lhs)
5. orthogonal array latin hypercube sampling (oa_lhs)
6. Box-Behnken (box_behnken)
7. central composite design (central_composite)

These methods all generate point sets that may be used to drive a set of computer experiments. Note that all of the DACE methods generated randomized designs, except for Box-Behnken and Central composite which are classical designs. That is, the grid sampling will generate a randomized grid, not what one typically thinks of as a grid of uniformly spaced points over a rectangular grid. Similar, the orthogonal array is a randomized version of an orthogonal array: it does not generate discrete, fixed levels.

In addition to the selection of the method, there are keywords that affect the method outputs:

1. main_effects
2. quality_metrics
3. variance_based_decomp

And keywords that affect the sampling:

1. fixed_seed
2. symbols
3. samples
4. seed
See Also

These keywords may also be of interest:

- fsu_cvt
- fsu_quasi_mc
- psuade_moat

7.2.69.1 grid

- Keywords Area
- method
- dace
- grid

Grid Sampling

Specification

**Alias**: none

**Argument(s)**: none

Description

The grid option in DACE will produce a randomized grid of points. If you are interested in a regular grid of points, use the multidimensional parameter study (under Parameter Studies) instead. Grid Sampling

7.2.69.2 random

- Keywords Area
- method
- dace
- random

Uses purely random Monte Carlo sampling to sample variables

Specification

**Alias**: none

**Argument(s)**: none
Description

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

```
method sampling
  sample_type random
  samples = 200
```

7.2.69.3 oas

- Keywords Area
- method
- dace
- oas

Orthogonal Array Sampling

Specification

Alias: none

Argument(s): none

Description

Orthogonal array sampling (OAS) is a widely used technique for running experiments and systematically testing factor effects. An orthogonal array sample can be described as a 4-tuple \((m;n;s;r)\), where \(m\) is the number of sample points, \(n\) is the number of input variables, \(s\) is the number of symbols, and \(r\) is the strength of the orthogonal array. The number of sample points, \(m\), must be a multiple of the number of symbols, \(s\). The number of symbols refers to the number of levels per input variable. The strength refers to the number of columns where we are guaranteed to see all the possibilities an equal number of times. Note that the DACE OAS capability produces a randomized orthogonal array: the samples for a particular level are randomized within that level.

If one examines the sample sets in an orthogonal array by looking at the rows as individual samples and columns as the variables sampled, one sees that the columns are orthogonal to each other in an orthogonal array. This feature is important in main effects analysis, which is a sensitivity analysis technique that identifies which variables have the most influence on the output.
Uses Latin Hypercube Sampling (LHS) to sample variables

Specification

Alias: none
Argument(s): none

Description

The *lhs* keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

Default Behavior

Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the *lhs* keyword must appear in conjunction with the *sample_type* keyword.

Usage Tips

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

Examples

```ini
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.69.5  **oa_lhs**

- Keywords Area
- method
- dace
- *oa_lhs*

Orthogonal Array Latin Hypercube Sampling

Specification

Alias: none
Argument(s): none
7.2 method

Description

The Orthogonal Array Latin Hypercube Sampling option in DACE produces a "latinized" version of an orthogonal array. That is, after the orthogonal array is generated, the samples go through a stratification process to produce samples that have been both orthogonalized and stratified.

See Also

These keywords may also be of interest:

- oas

7.2.69.6 box_behnken

- Keywords Area
- method
- dace
- box_behnken

Box-Behnken Design

Specification

Alias: none
Argument(s): none

Description

The Box-Behnken design is similar to a Central Composite design, with some differences. The Box-Behnken design is a quadratic design in that it does not contain an embedded factorial or fractional factorial design. In this design the treatment combinations are at the midpoints of edges of the process space and at the center, as compared with CCD designs where the extra points are placed at 'star points' on a circle outside of the process space. Box-Behken designs are rotatable (or near rotatable) and require 3 levels of each factor.

7.2.69.7 central_composite

- Keywords Area
- method
- dace
- central_composite

Central Composite Design

Specification

Alias: none
Argument(s): none
Description

A central composite design (CCD), contains an embedded factorial or fractional factorial design with a center points that is augmented with a group of “star points” that allow estimation of curvature.

Examples

See the User’s Manual for an example.

7.2.69.8 samples

- Keywords Area
- method
- dace
- samples

Number of samples for sampling-based methods

Specification

Alias: none
Argument(s): INTEGER
Default: 0

Description

The samples keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least dim+1 samples should be used, where “dim” is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least (dim+1)(dim+2)/2 samples are needed. For uncertainty quantification, we recommend at least 10*dim samples. For variance_based_decomp, we recommend hundreds to thousands of samples. Note that for variance_based_decomp, the number of simulations performed will be N*(dim+2).

Examples

method sampling
    sample_type lhs
    samples = 20

7.2.69.9 seed

- Keywords Area
- method
7.2 method

- dace
- seed

Seed of the random number generator

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```
method sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.69.10 fixed_seed

- Keywords Area
- method
- dace
- fixed_seed

Reuses the same seed value for multiple random sampling sets

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** not fixed; pattern varies run-to-run
Description

The `fixed_seed` flag is relevant if multiple sampling sets will be generated over the coarse of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the `seed` must also be specified.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed
```

7.2.69.11 main_effects

- Keywords Area
- method
- dace
- main_effects

ANOVA

Specification

Alias: none
Argument(s): none
Default: No main_effects

Description

The `main_effects` control prints Analysis-of-Variance main effects results (e.g., ANOVA tables with p-values per variable). The `main_effects` control is only operational with the orthogonal arrays or Latin Hypercube designs, not for Box Behnken or Central Composite designs.

Main effects is a sensitivity analysis method which identifies the input variables that have the most influence on the output. In main effects, the idea is to look at the mean of the response function when variable A (for example) is at level 1 vs. when variable A is at level 2 or level 3. If these mean responses of the output are statistically significantly different at different levels of variable A, this is an indication that variable A has a significant effect on the response. The orthogonality of the columns is critical in performing main effects analysis, since the column orthogonality means that the effects of the other variables “cancel out” when looking at the overall effect from one variable at its different levels.
Calculate metrics to assess the quality of quasi-Monte Carlo samples

Topics

This keyword is related to the topics:

- package_fsudace

Specification

Alias: none
Argument(s): none
Default: No quality_metrics

Description

quality_metrics calculates four quality metrics relating to the volumetric spacing of the samples. The four quality metrics measure different aspects relating to the uniformity of point samples in hypercubes. Desirable properties of such point samples are:

- are the points equally spaced
- do the points cover the region
- and are they isotropically distributed
- with no directional bias in the spacing

The four quality metrics we report are:

- $h$: the point distribution norm, which is a measure of uniformity of the point distribution
- $\chi$: a regularity measure, and provides a measure of local uniformity of a set of points
- $\tau$: the second moment trace measure
- $d$: the second moment determinant measure

All of these values are scaled so that smaller is better (the smaller the metric, the better the uniformity of the point distribution).

Examples

Complete explanation of these measures can be found in [39].
7.2.69.13 variance_based_decomp

- Keywords Area
- method
- dace
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into contributions from variables

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>drop_tolerance</td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate sensitivity indices through variance based decomposition using the keyword `variance_based_decomp`. These indicate how important the uncertainty in each input variable is in contributing to the output variance.

**Default Behavior**

Because of the computational cost, `variance_based_decomp` is turned off as a default.

If the user specified a number of samples, N, and a number of nondeterministic variables, M, variance-based decomposition requires the evaluation of $N \times (M+2)$ samples. **Note that specifying this keyword will increase the number of function evaluations above the number requested with the samples keyword since replicated sets of sample values are evaluated.**

**Expected Outputs**

When `variance_based_decomp` is specified, sensitivity indices for main effects and total effects will be reported. Main effects (roughly) represent the percent contribution of each individual variable to the variance in the model response. Total effects represent the percent contribution of each individual variable in combination with all other variables to the variance in the model response.

**Usage Tips**

To obtain sensitivity indices that are reasonably accurate, we recommend that N, the number of samples, be at least one hundred and preferably several hundred or thousands.

**Examples**

```verbatim
method,
sampling
  sample_type lhs
```
Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of [76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in [76] and [90].

7.2.69.13.1 drop_tolerance

- Keywords Area
- method
- dace
- variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance

Specification

Alias: none

Argument(s): REAL

Default: All VBD indices displayed

Description

The drop_tolerance keyword allows the user to specify a value below which sensitivity indices generated by variance_based_decomp are not displayed.

Default Behavior

By default, all sensitivity indices generated by variance_based_decomp are displayed.

Usage Tips

For polynomial_chaos, which outputs main, interaction, and total effects by default, the univariate_effects may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

```dakota
method,  
  sampling  
    sample_type lhs  
    samples = 100  
    variance_based_decomp  
    drop_tolerance = 0.001
```
7.2.69.14 symbols

- Keywords Area
- method
- dace
- symbols

Number of replications in the sample set

Specification

Alias: none
Argument(s): INTEGER
Default: default for sampling algorithm

Description

symbols is related to the number of levels per variable in the sample set (a larger number of symbols equates to more stratification and fewer replications). For example, if symbols = 7, each variable would be divided into seven levels.

7.2.69.15 model_pointer

- Keywords Area
- method
- dace
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none
Argument(s): STRING
Default: method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```xml
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians
```
7.2.70 fsu_cvt

- Keywords Area
- method
- fsu_cvt

Design of Computer Experiments - Centroidal Voronoi Tessellation

Topics

This keyword is related to the topics:

- package_fsudace
- design_and_analysis_of_computer_experiments

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>samples</td>
<td></td>
<td>Number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>fixed_seed</td>
<td></td>
<td>Reuses the same seed value for multiple random sampling sets</td>
</tr>
<tr>
<td>Optional</td>
<td>latinize</td>
<td></td>
<td>Adjust samples to improve the discrepancy of the marginal distributions</td>
</tr>
<tr>
<td>Optional</td>
<td>quality_metrics</td>
<td></td>
<td>Calculate metrics to assess the quality of quasi-Monte Carlo samples</td>
</tr>
</tbody>
</table>
### Description

The FSU Centroidal Voronoi Tessellation method (*fsu_cvt*) produces a set of sample points that are (approximately) a Centroidal Voronoi Tessellation. The primary feature of such a set of points is that they have good volumetric spacing; the points tend to arrange themselves in a pattern of cells that are roughly the same shape.

To produce this set of points, an almost arbitrary set of initial points is chosen, and then an internal set of iterations is carried out. These iterations repeatedly replace the current set of sample points by an estimate of the centroids of the corresponding Voronoi subregions. [18].

The user may generally ignore the details of this internal iteration. If control is desired, however, there are a few variables with which the user can influence the iteration. The user may specify:

- **max_iterations**, the number of iterations carried out
- **num_trials**, the number of secondary sample points generated to adjust the location of the primary sample points
- **trial_type**, which controls how these secondary sample points are generated

This method generates sets of uniform random variables on the interval [0, 1]. If the user specifies lower and upper bounds for a variable, the [0, 1] samples are mapped to the [lower, upper] interval.

### Theory

This method is designed to generate samples with the goal of low discrepancy. Discrepancy refers to the nonuniformity of the sample points within the hypercube.

Discrepancy is defined as the difference between the actual number and the expected number of points one would expect in a particular set B (such as a hyper-rectangle within the unit hypercube), maximized over all such sets. Low discrepancy sequences tend to cover the unit hypercube reasonably uniformly.
Centroidal Voronoi Tessellation does very well volumetrically: it spaces the points fairly equally throughout the space, so that the points cover the region and are isotropically distributed with no directional bias in the point placement. There are various measures of volumetric uniformity which take into account the distances between pairs of points, regularity measures, etc. Note that Centroidal Voronoi Tessellation does not produce low-discrepancy sequences in lower dimensions. The lower-dimension (such as 1-D) projections of Centroidal Voronoi Tessellation can have high discrepancy.

See Also

These keywords may also be of interest:

- dace
- fsu_quasi_mc
- psuade_moat

7.2.70.1 samples

- Keywords Area
- method
- fsu_cvt
- samples

Number of samples for sampling-based methods

Specification

Alias: none
Argument(s): INTEGER
Default: 0

Description

The samples keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least dim+1 samples should be used, where "dim" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \((\text{dim}+1)(\text{dim}+2)/2\) samples are needed. For uncertainty quantification, we recommend at least 10\(	ext{dim}\) samples. For variance_based_decomp, we recommend hundreds to thousands of samples. Note that for variance_based_decomp, the number of simulations performed will be \(N=\text{dim}+2\).

Examples

```
method
  sampling
    sample_type lhs
    samples = 20
```
7.2.70.2 seed

- Keywords Area
- method
- fsu_cvt
- seed

Seed of the random number generator

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**
If not specified, the seed is randomly generated.

**Expected Output**
If seed is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**
If a stochastic study was run without seed specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.70.3 fixed_seed

- Keywords Area
- method
- fsu_cvt
- fixed_seed

Reuses the same seed value for multiple random sampling sets
Keywords Area

Specification

Alias: none
Argument(s): none
Default: not fixed; pattern varies run-to-run

Description

The fixed_seed flag is relevant if multiple sampling sets will be generated over the course of a Dakota analysis. This occurs when using advance methods (e.g., surrogate-based optimization, optimization under uncertainty). The same seed value is reused for each of these multiple sampling sets, which can be important for reducing variability in the sampling results.

Default Behavior

The default behavior is to not use a fixed seed, as the repetition of the same sampling pattern can result in a modeling weakness that an optimizer could potentially exploit (resulting in actual reliabilities that are lower than the estimated reliabilities). For repeatable studies, the seed must also be specified.

Examples

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    fixed_seed

7.2.70.4 latinize
```

Adjust samples to improve the discrepancy of the marginal distributions

Specification

Alias: none
Argument(s): none
Default: No latinization

Description

The latinize control takes the samples and "latinizes" them, meaning that each original sample is moved so that it falls into one strata or bin in each dimension as in Latin Hypercube sampling. The default setting is NOT to latinize. However, one may be interested in doing this in situations where one wants better discrepancy of the 1-dimensional projections (the marginal distributions).
7.2.70.5 quality_metrics

- Keywords Area
- method
- fsu_cvt
- quality_metrics

Calculate metrics to assess the quality of quasi-Monte Carlo samples

Topics

This keyword is related to the topics:

- package_fсудace

Specification

Alias: none
Argument(s): none
Default: No quality_metrics

Description

quality_metrics calculates four quality metrics relating to the volumetric spacing of the samples. The four quality metrics measure different aspects relating to the uniformity of point samples in hypercubes. Desirable properties of such point samples are:

- are the points equally spaced
- do the points cover the region
- and are they isotropically distributed
- with no directional bias in the spacing

The four quality metrics we report are:

- h: the point distribution norm, which is a measure of uniformity of the point distribution
- chi: a regularity measure, and provides a measure of local uniformity of a set of points
- tau: the second moment trace measure
- d: the second moment determinant measure

All of these values are scaled so that smaller is better (the smaller the metric, the better the uniformity of the point distribution).

Examples

Complete explanation of these measures can be found in [39].
7.2.70.6 variance_based_decomp

- Keywords Area
- method
- fsu_cvt
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into contributions from variables

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no variance-based decomposition

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>drop_tolerance</td>
<td></td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

**Description**

Dakota can calculate sensitivity indices through variance based decomposition using the keyword `variance_based_decomp`. These indicate how important the uncertainty in each input variable is in contributing to the output variance.

**Default Behavior**

Because of the computational cost, `variance_based_decomp` is turned off as a default.

If the user specified a number of samples, N, and a number of nondeterministic variables, M, variance-based decomposition requires the evaluation of \( N \times (M+2) \) samples. **Note that specifying this keyword will increase the number of function evaluations above the number requested with the `samples` keyword since replicated sets of sample values are evaluated.**

**Expected Outputs**

When `variance_based_decomp` is specified, sensitivity indices for main effects and total effects will be reported. Main effects (roughly) represent the percent contribution of each individual variable to the variance in the model response. Total effects represent the percent contribution of each individual variable in combination with all other variables to the variance in the model response.

**Usage Tips**

To obtain sensitivity indices that are reasonably accurate, we recommend that N, the number of samples, be at least one hundred and preferably several hundred or thousands.

**Examples**

```
method,
  sampling
    sample_type lhs
```
Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of [76]: "The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input."

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in [76] and [90].

7.2.70.6.1 drop_tolerance

- Keywords Area
  - method
  - fsu_cvt
  - variance_based_decomp
  - drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance

Specification

Alias: none

Argument(s): REAL

Default: All VBD indices displayed

Description

The `drop_tolerance` keyword allows the user to specify a value below which sensitivity indices generated by `variance_based_decomp` are not displayed.

Default Behavior

By default, all sensitivity indices generated by `variance_based_decomp` are displayed.

Usage Tips

For `polynomial_chaos`, which outputs main, interaction, and total effects by default, the `univariate_effects` may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).

Examples

```
method,
sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
  drop_tolerance = 0.001
```
7.2.70.7 trial_type

- Keywords Area
- method
- fsu_cvt
- trial_type

Specify how the trial samples are generated

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** random

**Child Keywords:**

<table>
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<tr>
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<th>Description of Group Trial Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>grid</td>
<td>Samples on a regular grid</td>
</tr>
<tr>
<td></td>
<td></td>
<td>halton</td>
<td>Generate samples from a Halton sequence</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The user has the option to specify the method by which the trials are created to adjust the centroids. The `trial_type` can be one of three types:

- **random**, where points are generated randomly
- **halton**, where points are generated according to the Halton sequence
- **grid**, where points are placed on a regular grid over the hyperspace.

7.2.70.7.1 grid

- Keywords Area
- method
- fsu_cvt
- trial_type
- grid

Samples on a regular grid
7.2 method

Specification

Alias: none
Argument(s): none

Description

Points are placed on a regular grid over the hyperspace.

See Also

These keywords may also be of interest:

• trial_type

7.2.70.7.2 halton

• Keywords Area
• method
• fsu_cvt
• trial_type
• halton

Generate samples from a Halton sequence

Topics

This keyword is related to the topics:

• package_fsudace

Specification

Alias: none
Argument(s): none

Description

The quasi-Monte Carlo sequences of Halton are deterministic sequences determined by a set of prime bases. These sequences generate random numbers with the goal of filling a unit hypercube uniformly.

Generally, we recommend that the user leave the default setting for the bases, which are the lowest primes. Thus, if one wants to generate a sample set for 3 random variables, the default bases used are 2, 3, and 5 in the Halton sequence. To give an example of how these sequences look, the Halton sequence in base 2 starts with points 0.5, 0.25, 0.75, 0.125, 0.625, etc. The first few points in a Halton base 3 sequence are 0.33333, 0.66667, 0.11111, 0.44444, 0.77777, etc. Notice that the Halton sequence tends to alternate back and forth, generating a point closer to zero then a point closer to one. An individual sequence is based on a radix inverse function defined on a prime base. The prime base determines how quickly the [0,1] interval is filled in.
Theory

For more information about these sequences, see [44], [45], and [1].

7.2.70.7.3 random

- Keywords Area
- method
- fsu_cvt
- trial_type
- random

Uses purely random Monte Carlo sampling to sample variables

Specification

Alias: none
Argument(s): none

Description

The random keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

Default Behavior

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the random keyword must be specified in conjunction with the sample_type keyword.

Usage Tips

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

Examples

method
  sampling
    sample_type random
    samples = 200

7.2.70.8 num_trials

- Keywords Area
- method
- fsu_cvt
- num_trials

The number of secondary sample points generated to adjust the location of the primary sample points
7.2 method

Specification

Alias: none
Argument(s): INTEGER
Default: 10000

Description

In general, the variable with the most influence on the quality of the final sample set is num_trials, which determines how well the Voronoi subregions are sampled. Generally, num_trials should be "large", certainly much bigger than the number of sample points being requested; a reasonable value might be 10,000, but values of 100,000 or 1 million are not unusual.

7.2.7.0.9 max_iterations

• Keywords Area
• method
• fsu_cvt
• max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations
7.2.70.10  model_pointer

• Keywords Area
• method
• fsu_cvt
• model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
       0.1 0.2 0.6
       0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
    id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.71 psuade_moat

• Keywords Area

• method

• psuade_moat

Morris One-at-a-Time

Topics

This keyword is related to the topics:

• package_psuade

• design_and_analysis_of_computer_experiments

Specification

Alias: none

Argument(s): none

Child Keywords:
Description

The Morris One-At-A-Time (MOAT) method, originally proposed by Morris [64], is a screening method, designed to explore a computational model to distinguish between input variables that have negligible, linear and additive, or nonlinear or interaction effects on the output. The computer experiments performed consist of individually randomized designs which vary one input factor at a time to create a sample of its elementary effects.

The number of samples (samples) must be a positive integer multiple of (number of continuous design variable + 1) and will be automatically adjusted if misspecified.

The number of partitions (partitions) applies to each variable being studied and must be odd (the number of MOAT levels per variable is partitions + 1). This will also be adjusted at runtime as necessary.

For information on practical use of this method, see [76].

Theory

With MOAT, each dimension of a $k$–dimensional input space is uniformly partitioned into $p$ levels, creating a grid of $p^k$ points $x \in \mathbb{R}^k$ at which evaluations of the model $y(x)$ might take place. An elementary effect corresponding to input $i$ is computed by a forward difference

$$d_i(x) = \frac{y(x + \Delta e_i) - y(x)}{\Delta},$$

where $e_i$ is the $i$th coordinate vector, and the step $\Delta$ is typically taken to be large (this is not intended to be a local derivative approximation). In the present implementation of MOAT, for an input variable scaled to $[0, 1]$, $\Delta = \frac{p}{2(p-1)}$, so the step used to find elementary effects is slightly larger than half the input range.

The distribution of elementary effects $d_i$ over the input space characterizes the effect of input $i$ on the output of interest. After generating $r$ samples from this distribution, their mean,

$$\mu_i = \frac{1}{r} \sum_{j=1}^{r} d_i^{(j)}$$

modified mean

$$\mu_i^* = \frac{1}{r} \sum_{j=1}^{r} |d_i^{(j)}|,$$

(using absolute value) and standard deviation

$$\sigma_i = \sqrt{\frac{1}{r} \sum_{j=1}^{r} \left( d_i^{(j)} - \mu_i \right)^2}$$
are computed for each input \( i \). The mean and modified mean give an indication of the overall effect of an input on the output. Standard deviation indicates nonlinear effects or interactions, since it is an indicator of elementary effects varying throughout the input space.

7.2.71.1 partitions

- Keywords Area
- method
- psuade_moat
- partitions

Number of partitions of each variable

Specification

Alias: none

Argument(s): INTEGERLIST

Default: 3

Description

Described on the parent page, psuade_moat

7.2.71.2 samples

- Keywords Area
- method
- psuade_moat
- samples

Number of samples for sampling-based methods

Specification

Alias: none

Argument(s): INTEGER

Default: 0

Description

The samples keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips
To obtain linear sensitivities or to construct a linear response surface, at least $\text{dim}+1$ samples should be used, where "dim" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least $(\text{dim}+1)(\text{dim}+2)/2$ samples are needed. For uncertainty quantification, we recommend at least $10\times\text{dim}$ samples. For \texttt{variance\_based\_decomp}, we recommend hundreds to thousands of samples. Note that for \texttt{variance\_based\_decomp}, the number of simulations performed will be $N\times(\text{dim}+2)$.

### Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 20

7.2.71.3 seed

- Keywords Area
- method
- psuade_moat
- seed

Seed of the random number generator

### Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

### Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

### Default Behavior

If not specified, the seed is randomly generated.

### Expected Output

If \texttt{seed} is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

### Usage Tips

If a stochastic study was run without \texttt{seed} specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

### Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```
7.2 method

7.2.71.4 model_pointer

- Keywords Area
- method
- psuade_moat
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:
- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The **model_pointer** is used to specify which **model** block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a **model** block in the Dakota input file that has a corresponding **id_model** with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a **model_pointer** for each method is imperative.

See **block_pointer** for details about pointers.

**Examples**

```plaintext
environment
tabular_data
method_pointer = 'UQ'

method
id_method = 'UQ'
model_pointer = 'SURR'
sampling,
samples = 10
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
0.1 0.2 0.6
0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
```
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0..
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.72 local_evidence

- Keywords Area
- method
- local_evidence

Evidence theory with evidence measures computed with local optimization methods

Topics

This keyword is related to the topics:

- epistemic_uncertainty_quantification_methods
- evidence_theory

Specification

Alias: nond_local_evidence
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Optimization Solver (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>sqp</td>
<td>Uses a sequential quadratic programming method for underlying optimization</td>
<td></td>
</tr>
<tr>
<td></td>
<td>nip</td>
<td>Uses a nonlinear interior point method for underlying optimization</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Two local optimization methods are available: `sqp` (sequential quadratic programming) or `nip` (nonlinear interior point method).

**Additional Resources**

See the topic page `evidence_theory` for important background information and usage notes.

Refer to `variable_support` for information on supported variable types.

**See Also**

These keywords may also be of interest:

- `global_evidence`
- `global_interval_est`
Uses a sequential quadratic programming method for underlying optimization

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The *sqp* keyword directs Dakota to use a sequential quadratic programming method to solve that problem. A sequential quadratic programming solves a sequence of linearly constrained quadratic optimization problems to arrive at the solution to the optimization problem.

Uses a nonlinear interior point method for underlying optimization

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The *nip* keyword directs Dakota to use a nonlinear interior point to solve that problem. A nonlinear interior point method traverses the interior of the feasible region to arrive at the solution to the optimization problem.
7.2.7.2.3  response_levels

- Keywords Area
- method
- local_evidence
- response_levels

Values at which to estimate desired statistics for each response

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_response_levels</td>
<td></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td>compute</td>
<td></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**

If `response_levels` are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If `response_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The `num_response_levels` is used to specify which arguments of the `response_level` correspond to which response.
Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num_response_levels key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

```dakota
method,
  sampling,
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                       6.0e+04 6.5e+04 7.0e+04
                       3.5e+05 4.0e+05 4.5e+05
variables,
  normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
    num_pairs = 3 4
    abscissas = 5 8 10 .1 .2 .3 .4
    counts = 17 21 0 12 24 12 0
    descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
  system asynch evaluationConcurrency = 5
  analysis_driver = 'text_book'
responses,
  response_functions = 3
  no_gradients
  no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
</tbody>
</table>
3.6000000000e+11 4.0000000000e+11 4.2500000000e-12
4.0000000000e+11 4.4000000000e+11 3.7500000000e-12
4.4000000000e+11 5.4196114379e+11 2.2557612778e-12

PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:
Complementary Cumulative Distribution Function (CCDF) for response_fn_1:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.

7.2.72.3.1 num_response_levels

- Keywords Area
- method
- local_evidence
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST
**Default:** response_levels evenly distributed among response functions

**Description**

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

**Default Behavior**

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

**Expected Outputs**

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

**Examples**

```
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.7 2.3.2 compute

- Keywords Area
- method
- local_evidence
- response_levels
- compute

Selection of statistics to compute at each response level

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
</tbody>
</table>
### Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

#### Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

#### Expected Output

The type of statistics specified by `compute` will be reported for each response level.

#### Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach. CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

#### Examples

```plaintext
method sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                    6.0e+04 6.5e+04 7.0e+04
                    3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```

#### 7.2.72.3.2.1 probabilities

- Keywords Area
- method
- local_evidence
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels.
The *probabilities* keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If *response_levels* is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the *probabilities* keyword should be specified in conjunction with the *compute* keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

**Examples**

```xml
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
  compute probabilities
```

7.2.72.3.2.2 *gen_reliabilities*

- *Keywords Area*
- *method*
- *local_evidence*
- *response_levels*
- *compute*
- *gen_reliabilities*

Computes generalized reliabilities associated with response levels

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The *gen_reliabilities* keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.
Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary_distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                     6.0e+04 6.5e+04 7.0e+04
                     3.5e+05 4.0e+05 4.5e+05
    compute gen_reliabilities
```

### 7.2.72.3.2.3 system

- **Keywords Area**
- **method**
- **local_evidence**
- **response_levels**
- **compute**
- **system**

Compute system reliability (series or parallel)

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group System Reliability Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td></td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
<tr>
<td></td>
<td></td>
<td>parallel</td>
<td>Aggregate response statistics assuming a parallel system</td>
</tr>
</tbody>
</table>

**Description**

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.
For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.72.3.2.4 series

- Keywords Area
- method
- local_evidence
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.

7.2.72.3.2.5 parallel

- Keywords Area
- method
- local_evidence
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

Alias: none
Argument(s): none

Description

See parent keyword system for description.
7.2.72.4 probability_levels

- Keywords Area
- method
- local_evidence
- probability_levels

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Examples**

The Dakota input file below specifies a sampling method with probability levels of interest.

```plaintext
method,
sampling,
samples = 100 seed = 1
complementary distribution
probability_levels = 1. 66.33 0.
1.8 .5 .0.
1.3 .2 .0.

variables,
normal_uncertain = 2
means = 248.89, 593.33
std_deviations = 12.4, 29.7
descriptors = 'TF1n' 'TF2n'
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
uniform_uncertain = 2
lower_bounds = 199.3, 474.63
upper_bounds = 298.5, 712.
descriptors = 'TF1u', 'TF2u'

weibull_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = 'TF1w', 'TF2w'

histogram_bin_uncertain = 2
num_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = 'TF1h', 'TF2h'

histogram_point_uncertain
real = 1
num_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = 'TF3h'

interface,
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses,
response_functions = 3
no_gradients
no_hessians

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

- PDF for response_fn_1:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.138477497e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.145412231e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.419614379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

- PDF for response_fn_2:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465755e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

- PDF for response_fn_3:
<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.699721413e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.699721413e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.127492534e-06</td>
</tr>
</tbody>
</table>

- Level mappings for each response function:
<p>| Complementary Cumulative Distribution Function (CCDF) for response_fn_1: |</p>
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td>2.7604749078e+11</td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.419614379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Complementary Cumulative Distribution Function (CCDF) for response_fn_2:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Complementary Cumulative Distribution Function (CCDF) for response_fn_3:
<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.72.4.1 num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.72.5 gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value
Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_gen_reliability_levels</td>
<td>gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.72.5.1 num_gen_reliability_levels

- Keywords Area
- method
- local_evidence
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** gen_reliability_levels evenly distributed among response functions
Description

See parent page

7.2.72.6 distribution

- Keywords Area
- method
- local_evidence
- distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group 1 (CDF/CCDF)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td></td>
<td>Distribution Type</td>
<td>cumulative</td>
<td>Computes statistics</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>according to cumulative</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>according to complementary</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>cumulative cumulative</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

method sampling
Computes statistics according to cumulative functions

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must be appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

Computes statistics according to complementary cumulative functions

```plaintext
```
7.2 method

Specification

**Alias:** none

**Argument(s):** none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

**Default Behavior**

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must appear in conjunction with the `distribution` keyword.

**Expected Outputs**

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary
```

7.2.7.2.7 model_pointer

- **Keywords Area**
- **method**
- **local_evidence**
- **model_pointer**

Identifier for model block to be used by a method

**Topics**

This keyword is related to the topics:

- **block_pointer**

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)
Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
        response_levels = 0.1 0.2 0.6
           0.1 0.2 0.6
           0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
    id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
    id_method = 'DACE'
    model_pointer = 'DACE_M'
    sampling sample_type lhs
        samples = 121 seed = 5034 rng rnum2

model
    id_model = 'DACE_M'
    single
    interface_pointer = 'I1'

variables
    uniform_uncertain = 2
        lower_bounds = 0.  0.
        upper_bounds = 1.  1.
        descriptors = 'x1' 'x2'

interface
    id_interface = 'I1'
    system asynch evaluation_concurrency = 5
    analysis_driver = 'text_book'

responses
    response_functions = 3
    no_gradients
    no_hessians
```
7.2.73 local_interval_est

Keywords Area

- method
- local_interval_est

Interval analysis using local optimization

Topics

This keyword is related to the topics:

- uncertainty_quantification
- epistemic_uncertainty_quantification_methods
- interval_estimation

Specification

Alias: nond_local_interval_est

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Optimization Solver (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>sqp</td>
<td>sqp</td>
<td>Uses a sequential quadratic programming method for underlying optimization</td>
</tr>
<tr>
<td></td>
<td>nip</td>
<td>nip</td>
<td>Uses a nonlinear interior point method for underlying optimization</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-tolerance</td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
</tbody>
</table>
Description

Interval analysis using local methods (local_interval_est). If the problem is amenable to local optimization methods (e.g. can provide derivatives or use finite difference method to calculate derivatives), then one can use one of two local methods to calculate these bounds.

- sqp
- nip

Additional Resources

Refer to variable_support for information on supported variable types.

Theory

In interval analysis, one assumes that nothing is known about an epistemic uncertain variable except that its value lies somewhere within an interval. In this situation, it is NOT assumed that the value has a uniform probability of occurring within the interval. Instead, the interpretation is that any value within the interval is a possible value or a potential realization of that variable. In interval analysis, the uncertainty quantification problem is one of determining the resulting bounds on the output (defining the output interval) given interval bounds on the inputs. Again, any output response that falls within the output interval is a possible output with no frequency information assigned to it.

See Also

These keywords may also be of interest:

- global_evidence
- global_interval_est
- local_evidence

7.2.73.1 sqp

- Keywords Area
- method
- local_interval_est
- sqp

Uses a sequential quadratic programming method for underlying optimization

Specification

**Alias:** none

**Argument(s):** none
Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The `sqp` keyword directs Dakota to use a sequential quadratic programming method to solve that problem. A sequential quadratic programming solves a sequence of linearly constrained quadratic optimization problems to arrive at the solution to the optimization problem.

7.2.73.2 nip

- Keywords Area
- method
- local_interval_est
- nip

Uses a nonlinear interior point method for underlying optimization

Specification

Alias: none
Argument(s): none

Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The `nip` keyword directs Dakota to use a nonlinear interior point to solve that problem. A nonlinear interior point method traverses the interior of the feasible region to arrive at the solution to the optimization problem.

7.2.73.3 convergence_tolerance

- Keywords Area
- method
- local_interval_est
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4
Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by `convergence_tolerance`, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See `nl2sol`
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.73.4 `model_pointer`

- Keywords Area
- method
- local_interval_est
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.
Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = ’UQ’

method
  id_method = ’UQ’
  model_pointer = ’SURR’
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
    0.1 0.2 0.6
    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative
model
  id_model = ’SURR’
  surrogate global,
  dace_method_pointer = ’DACE’
  polynomial quadratic
method
  id_method = ’DACE’
  model_pointer = ’DACE_M’
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2
model
  id_model = ’DACE_M’
  single
  interface_pointer = ’I1’
variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = ’x1’ ’x2’
interface
  id_interface = ’I1’
  system asynch evaluation_concurrency = 5
  analysis_driver = ’text_book’
responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.74  local_reliability

- Keywords Area
- method
- local_reliability
Local reliability method

Topics

This keyword is related to the topics:

- uncertainty_quantification
- reliability_methods

Specification

Alias: nond_local_reliability

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>mpp_search</td>
<td>Specify which MPP search option to use</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>response_levels</td>
<td>Values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>probability_levels</td>
<td>Specify probability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>reliability_levels</td>
<td>Specify reliability levels at which the response values will be estimated</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>gen_reliability_levels</td>
<td>Specify generalized reliability levels at which to estimate the corresponding response value</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>distribution</td>
<td>Selection of cumulative or complementary cumulative functions</td>
</tr>
</tbody>
</table>
Description

Local reliability methods compute approximate response function distribution statistics based on specified uncertain variable probability distributions. Each of the local reliability methods can compute forward and inverse mappings involving response, probability, reliability, and generalized reliability levels.

The forward reliability analysis algorithm of computing reliabilities/probabilities for specified response levels is called the Reliability Index Approach (RIA), and the inverse reliability analysis algorithm of computing response levels for specified probability levels is called the Performance Measure Approach (PMA).

The different RIA/PMA algorithm options are specified using the mpp_search specification which selects among different limit state approximations that can be used to reduce computational expense during the MPP searches.

Theory

The Mean Value method (MV, also known as MVFOSM in [43]) is the simplest, least-expensive method in that it estimates the response means, response standard deviations, and all CDF/CCDF forward/inverse mappings from a single evaluation of response functions and gradients at the uncertain variable means. This approximation can have acceptable accuracy when the response functions are nearly linear and their distributions are approximately Gaussian, but can have poor accuracy in other situations.

All other reliability methods perform an internal nonlinear optimization to compute a most probable point (MPP) of failure. A sign convention and the distance of the MPP from the origin in the transformed standard normal space (“u-space”) define the reliability index, as explained in the section on Reliability Methods in the Uncertainty Quantification chapter of the Users Manual [5]. Also refer to variable_support for additional information on supported variable types for transformations to standard normal space. The reliability can then be converted to a probability using either first- or second-order integration, may then be refined using importance sampling, and finally may be converted to a generalized reliability index.

See Also

These keywords may also be of interest:

- adaptive_sampling
7.2.74.1 mpp_search

**Keywords Area**
- method
- local_reliability
- mpp_search

Specify which MPP search option to use

**Topics**
This keyword is related to the topics:
- uncertainty_quantification
- reliability_methods

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: No MPP search (MV method)

**Child Keywords**:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Required (Choose One)</td>
<td>MPP</td>
<td>x_taylor_mean</td>
<td>Form Taylor series approximation in &quot;x-space&quot; at variable means</td>
</tr>
<tr>
<td></td>
<td>Approximation (Group 1)</td>
<td>u_taylor_mean</td>
<td>Form Taylor series approximation in &quot;u-space&quot; at variable means</td>
</tr>
<tr>
<td></td>
<td></td>
<td>x_taylor_mpp</td>
<td>X-space Taylor series approximation with iterative updates</td>
</tr>
</tbody>
</table>
### Description

The `x_taylor_mean` MPP search option performs a single Taylor series approximation in the space of the original uncertain variables ("x-space") centered at the uncertain variable means, searches for the MPP for each response/probability level using this approximation, and performs a validation response evaluation at each predicted MPP. This option is commonly known as the Advanced Mean Value (AMV) method. The `u_taylor_mean` option is identical to the `x_taylor_mean` option, except that the approximation is performed in u-space. The `x_taylor_mpp` approach starts with an x-space Taylor series at the uncertain variable means, but iteratively updates the Taylor series approximation at each MPP prediction until the MPP converges. This option is commonly...
known as the AMV+ method. The \texttt{u_taylor_mpp} option is identical to the \texttt{x_taylor_mpp} option, except that all approximations are performed in \textit{u-space}. The order of the Taylor-series approximation is determined by the corresponding \texttt{responses} specification and may be first or second-order. If second-order (methods named \texttt{AMV^2} and \texttt{AMV^2+} in [22]), the series may employ analytic, finite difference, or quasi Hessians (BFGS or SR1).

The \texttt{x\_two\_point} MPP search option uses an \textit{x-space} Taylor series approximation at the uncertain variable means for the initial MPP prediction, then utilizes the Two-point Adaptive Nonlinear Approximation (TANA) outlined in [95] for all subsequent MPP predictions. The \texttt{u\_two\_point} approach is identical to \texttt{x\_two\_point}, but all the approximations are performed in \textit{u-space}. The \texttt{x\_taylor\_mpp} and \texttt{u\_taylor\_mpp}, \texttt{x\_two\_point} and \texttt{u\_two\_point} approaches utilize the \texttt{max\_iterations} and \texttt{convergence\_tolerance} method independent controls to control the convergence of the MPP iterations (the maximum number of MPP iterations per level is limited by \texttt{max\_iterations}, and the MPP iterations are considered converged when \( \| \mathbf{u}(k+1) - \mathbf{u}(k) \|_2 \leq \text{convergence\_tolerance} \)). And, finally, the \texttt{no\_approx} option performs the MPP search on the original response functions without the use of any approximations. The optimization algorithm used to perform these MPP searches can be selected to be either sequential quadratic programming (uses the \texttt{npsol\_sqp} optimizer) or nonlinear interior point (uses the \texttt{optpp\_q\_newton} optimizer) algorithms using the \texttt{sqp} or \texttt{nip} keywords.

In addition to the MPP search specifications, one may select among different integration approaches for computing probabilities at the MPP by using the \texttt{integration} keyword followed by either \texttt{first\_order} or \texttt{second\_order}. Second-order integration employs the formulation of [51] (the approach of [12] and the correction of [52] are also implemented, but are not active). Combining the \texttt{no\_approx} option of the MPP search with first- and second-order integrations results in the traditional first- and second-order reliability methods (FORM and SORM). These integration approximations may be subsequently refined using importance sampling. The refinement specification allows the selection of basic importance sampling (\texttt{import}), adaptive importance sampling (\texttt{adapt\_import}), or multimodal adaptive importance sampling (\texttt{mm\_adapt\_import}), along with the specification of number of samples (\texttt{samples}) and random seed (\texttt{seed}). Additional details on these methods are available in [24] and [22] and in the Uncertainty Quantification Capabilities chapter of the Users Manual [5].

7.2.74.1.1 \texttt{x\_taylor\_mean}

- Keywords Area
- method
- local\_reliability
- mpp\_search
- \texttt{x\_taylor\_mean}

Form Taylor series approximation in "x-space" at variable means

Topics

This keyword is related to the topics:

- reliability\_methods

Specification

\textbf{Alias:} none

\textbf{Argument(s):} none

Description

This \texttt{mpp\_search} option performs a single Taylor series approximation in the space of the original uncertain variables ("x-space") centered at the uncertain variable means, searches for the MPP for each response/probability level using this approximation, and performs a validation response evaluation at each predicted MPP. This option is commonly known as the Advanced Mean Value (AMV) method.
7.2.74.1.2  u_taylor_mean

- Keywords Area
- method
- local_reliability
- mpp_search
- u_taylor_mean

Form Taylor series approximation in "u-space" at variable means

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: none
Argument(s): none

Description

This mpp_search option performs a single Taylor series approximation in the transformed space of the uncertain variables ("u-space") centered at the uncertain variable means. This option is commonly known as the Advanced Mean Value (AMV) method, but is performed in u-space instead of x-space.

7.2.74.1.3  x_taylor_mpp

- Keywords Area
- method
- local_reliability
- mpp_search
- x_taylor_mpp

X-space Taylor series approximation with iterative updates

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: none
Argument(s): none
This mpp_search option starts with an x-space Taylor series at the uncertain variable means, but iteratively updates the Taylor series approximation at each MPP prediction until the MPP converges. This option is commonly known as the AMV+ method.

7.2.74.1.4  u_taylor_mpp

- Keywords Area
- method
- local_reliability
- mpp_search
- u_taylor_mpp

U-space Taylor series approximation with iterative updates

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: none

Argument(s): none

Description

This mpp_search option starts with a u-space Taylor series at the uncertain variable means, and iteratively updates the Taylor series approximation at each MPP prediction until the MPP converges. This option is commonly known as the AMV+ method and is identify to x_taylor_mpp except that it is performed in u-space.

7.2.74.1.5  x_two_point

- Keywords Area
- method
- local_reliability
- mpp_search
- x_two_point

Predict MPP using Two-point Adaptive Nonlinear Approximation in "x-space"

Topics

This keyword is related to the topics:

- reliability_methods
7.2 method 3153

Specification

Alias: none

Argument(s): none

Description

This mpp_search option uses an x-space Taylor series approximation at the uncertain variable means for the initial MPP prediction, then utilizes the Two-point Adaptive Nonlinear Approximation (TANA) outlined in [ Xu98 "Xu and Grandhi, 1998"] for all subsequent MPP predictions.

7.2.74.1.6 u_two_point

- Keywords Area
- method
- local_reliability
- mpp_search
- u_two_point

Predict MPP using Two-point Adaptive Nonlinear Approximation in "u-space"

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: none

Argument(s): none

Description

This mpp_search option is identical to x_two_point, but it performs the Two-point Adaptive Nonlinear Approximation (TANA) in u-space instead of x-space.

7.2.74.1.7 x_multi_point

- Keywords Area
- method
- local_reliability
- mpp_search
- x_multi_point

MPP search for local reliability based on QMEA multi-point approximation in u-space
Local reliability methods that search for the most probable point of failure (MPP) may use a surrogate-based optimization procedure. The \textit{x}\textunderscore multi\_point surrogate option creates a QMEA approximation in the original random variable space ("x-space") from accumulating data from multiple expansion points.

This capability is \textbf{experimental}.

7.2.74.1.8 \texttt{u\_multi\_point}

- **Keywords Area**
- **method**
- **local\_reliability**
- **mpp\_search**
- **u\_multi\_point**

MPP search for local reliability based on QMEA multi-point approximation in x-space

**Specification**

\textbf{Alias}: none

\textbf{Argument(s)}: none

**Description**

Local reliability methods that search for the most probable point of failure (MPP) may use a surrogate-based optimization procedure. The \texttt{u}\textunderscore multi\_point surrogate option creates a QMEA approximation in the transformed probability space ("u-space") from accumulating data from multiple expansion points.

This capability is \textbf{experimental}.

7.2.74.1.9 \texttt{no\_approx}

- **Keywords Area**
- **method**
- **local\_reliability**
- **mpp\_search**
- **no\_approx**

Perform MPP search on original response functions (use no approximation)

**Topics**

This keyword is related to the topics:

- **reliability\_methods**
7.2 method

Specification

Alias: none
Argument(s): none

Description

This mpp_search option performs the MPP search on the original response functions without the use of any approximations. Note that the use of the no_approx MPP search with first-order probability integration results in the traditional reliability method called FORM (First-Order Reliability Method). Similarly, the use of no_approx with second-order probability integration results in SORM (Second-Order Reliability Method).

7.2.74.1.10 sqp

- Keywords Area
- method
- local_reliability
- mpp_search
- sqp

Uses a sequential quadratic programming method for underlying optimization

Specification

Alias: none
Argument(s): none
Default: sqp

Description

Many uncertainty quantification methods solve a constrained optimization problem under the hood. The sqp keyword directs Dakota to use a sequential quadratic programming method to solve that problem. A sequential quadratic programming solves a sequence of linearly constrained quadratic optimization problems to arrive at the solution to the optimization problem.

7.2.74.1.11 nip

- Keywords Area
- method
- local_reliability
- mpp_search
- nip

Uses a nonlinear interior point method for underlying optimization
Many uncertainty quantification methods solve a constrained optimization problem under the hood. The \texttt{nip} keyword directs Dakota to use a nonlinear interior point to solve that problem. A nonlinear interior point method traverses the interior of the feasible region to arrive at the solution to the optimization problem.

### Integration approach

This keyword is related to the topics:

- reliability\_methods

### Specification

**Alias:** none  
**Argument(s):** none  
**Default:** sqp
Description

This keyword controls how the probabilities at the MPP are computed: integration is followed by either first_order or second_order, indicating the order of the probability integration.

7.2.74.1.12.1 first_order

- Keywords Area
- method
- local_reliability
- mpp_search
- integration
- first_order

First-order integration scheme

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: none

Argument(s): none

Description

First-order integration in local reliability methods uses the minimum Euclidean distance from the origin to the most probable point (MPP) in transformed space to compute the probability of failure. This distance, commonly called the reliability index Beta, is used to calculate the probability of failure by calculating the standard normal cumulative distribution function at -Beta.

7.2.74.1.12.2 second_order

- Keywords Area
- method
- local_reliability
- mpp_search
- integration
- second_order

Second-order integration scheme

Topics

This keyword is related to the topics:

- reliability_methods
Specification

Alias: none
Argument(s): none

Description

Second-order integration in local reliability methods modifies the first-order integration approach to apply a curvature correction. This correction is based on the formulation of [Hoh88 "Hohenbichler and Rackwitz, 1988"].

7.2.74.1.12.3 probability_refinement

• Keywords Area
• method
• local_reliability
• mpp_search
• integration
• probability_refinement

Allow refinement of probability and generalized reliability results using importance sampling

Topics

This keyword is related to the topics:

• reliability_methods

Specification

Alias: sample_refinement
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Required (Choose One)</th>
<th>Description of Group Importance Sampling Approach (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>mm_adapt_import</td>
<td>Importance sampling option for probability refinement</td>
</tr>
</tbody>
</table>
The `probability_refinement` allows refinement of probability and generalized reliability results using importance sampling. If one specifies `probability_refinement`, there are some additional options. One can specify which type of importance sampling to use (`import`, `adapt_import`, or `mm_adapt_import`). Additionally, one can specify the number of refinement samples to use with `refinement_samples` and the seed to use with `seed`.

The `probability_refinement` density reweighting accounts originally was developed based on Gaussian distributions. It now accounts for additional non-Gaussian cases.

### 7.2.74.1.12.4 import

- **Keywords** Area
- **method**
- **local_reliability**
- **mpp_search**
- **integration**
- **probability_refinement**
- **import**

Importance sampling option for probability refinement

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

`import` centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level).

### 7.2.74.1.12.5 adapt_import

- **Keywords** Area
- **method**
- **local_reliability**
- **mpp_search**
- **integration**
Importance sampling option for probability refinement

Specification

Alias: none

Argument(s): none

Description

adapt_import centers a sampling density at one of the initial LHS samples identified in the failure region. It then generates the importance samples, weights them by their probability of occurrence given the original density, and calculates the required probability (CDF or CCDF level). This continues iteratively until the failure probability estimate converges.

7.2.74.1.12.6 mm_adapt_import

Importance sampling option for probability refinement

Specification

Alias: none

Argument(s): none

Description

mm_adapt_import starts with all of the samples located in the failure region to build a multimodal sampling density. First, it uses a small number of samples around each of the initial samples in the failure region. Note that these samples are allocated to the different points based on their relative probabilities of occurrence: more probable points get more samples. This early part of the approach is done to search for “representative” points. Once these are located, the multimodal sampling density is set and then mm_adapt_import proceeds similarly to adapt_import (sample until convergence).
7.2 method

• mpp_search
• integration
• probability_refinement
• refinement_samples

Number of samples used to refine a probability estimate or sampling design.

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Probability estimate: Specify the (scalar) number of samples used to improve a probability estimate. If using uni-modal sampling all samples are assigned to the sampling center. If using multi-modal sampling the samples are split between multiple samples according to some internally computed weights.

Sampling design: Specify one or a sequence of refinement samples to augment the initial samples in a sampling design.

7.2.74.1.12.8 seed

• Keywords Area
• method
• local_reliability
• mpp_search
• integration
• probability_refinement
• seed

Seed of the random number generator

Specification

Alias: none
Argument(s): INTEGER
Default: system-generated (non-repeatable)

Description

The random seed control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

Default Behavior

If not specified, the seed is randomly generated.

Expected Output
If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 15347
```

7.2.74.2 `response_levels`

- **Keywords Area**
- **method**
- **local_reliability**
- **response_levels**

Values at which to estimate desired statistics for each response

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF probabilities/reliabilities to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td><code>num_response_levels</code></td>
<td>Number of values at which to estimate desired statistics for each response</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td><code>compute</code></td>
<td>Selection of statistics to compute at each response level</td>
</tr>
</tbody>
</table>

**Description**

The `response_levels` specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).

**Default Behavior**
If \texttt{response\_levels} are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

**Expected Outputs**

If \texttt{response\_levels} are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the \texttt{probability\_levels} and/or \texttt{response\_levels} in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Usage Tips**

The \texttt{num\_response\_levels} is used to specify which arguments of the \texttt{response\_level} correspond to which response.

**Examples**

For example, specifying a \texttt{response\_level} of 52.3 followed with \texttt{compute probabilities} will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

\begin{verbatim}
response\_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num\_response\_levels = 2 4 3
\end{verbatim}

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the \texttt{num\_response\_levels} key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

\begin{verbatim}
method,
sampling,
samples = 100 seed = 1
complementary distribution
response\_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
variables,
normal\_uncertain = 2
means = 248.89, 593.33
std\_deviations = 12.4, 29.7
descriptors = ’TF1n’, ’TF2n’
uniform\_uncertain = 2
lower\_bounds = 199.3, 474.63
upper\_bounds = 298.5, 712.
descriptors = ’TF1u’, ’TF2u’
weibull\_uncertain = 2
alphas = 12., 30.
betas = 250., 590.
descriptors = ’TF1w’, ’TF2w’
histogram\_bin\_uncertain = 2
num\_pairs = 3 4
abscissas = 5 8 10 .1 .2 .3 .4
counts = 17 21 0 12 24 12 0
descriptors = ’TF1h’, ’TF2h’
histogram\_point\_uncertain
real = 1
num\_pairs = 2
abscissas = 3 4
counts = 1 1
descriptors = ’TF3h’
\end{verbatim}
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.3601733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.5000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>3.8000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>2.3000000000e-01</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>2.9000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>5.2000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>9.0000000000e-02</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.5000000000e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.74.2.1 num_response_levels

- Keywords Area
- method
- local_reliability
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The num_response_levels keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If num_response_levels is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

method
sampling
  samples = 100
  seed = 34785
  num_response_levels = 1 1 1
  response_levels = 0.5 0.5 0.5

7.2.74.2.2 compute

- Keywords Area
- method
- local_reliability
- response_levels
- compute

Selection of statistics to compute at each response level
Specification

Alias: none
Argument(s): none
Default: probabilities
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>relabilities</td>
<td>Computes reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>system</td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

Description

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

Default Behavior

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

Expected Output

The type of statistics specified by `compute` will be reported for each response level.

Usage Tips

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

Examples

```plaintext
method
sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04
```
7.2 method

3.5e+05 4.0e+05 4.5e+05
compute reliabilities

7.2.74.2.2.1 probabilities

- Keywords Area
- method
- local_reliability
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

Specification

Alias: none
Argument(s): none

Description

The **probabilities** keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

Default Behavior

If **response_levels** is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the **probabilities** keyword should be specified in conjunction with the **compute** keyword.

Expected Outputs

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

Examples

```dakota
method
sampling
  sample_type random
  samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
  6.0e+04 6.5e+04 7.0e+04
  3.5e+05 4.0e+05 4.5e+05
compute probabilities
```

7.2.74.2.2.2 reliabilities

- Keywords Area
- method
- local_reliability
- response_levels
• compute

• reliabilities

Computes reliabilities associated with response levels

Specification

Alias: none

Argument(s): none

Description

The reliabilities keyword directs Dakota to compute reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If response_levels is specified, the reliabilities are not computed by default. To change this behavior, the reliabilities keyword should be specified in conjunction with the compute keyword.

Expected Outputs

The Dakota output is a set of response level-reliability pairs according to the distribution defined.

Examples

method
  sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
    6.0e+04 6.5e+04 7.0e+04
    3.5e+05 4.0e+05 4.5e+05
  compute reliabilities

7.2.74.2.2.3 gen_reliabilities

• Keywords Area

• method

• local_reliability

• response_levels

• compute

• gen_reliabilities

Computes generalized reliabilities associated with response levels

Specification

Alias: none

Argument(s): none
Description

The `gen_reliabilities` keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

Default Behavior

If `response_levels` is specified, the generalized reliabilities are not computed by default. To change this behavior, the `gen_reliabilities` keyword should be specified in conjunction with the `compute` keyword.

Expected Outputs

The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

Examples

```plaintext
method
  sampling
    sample_type random
    samples = 100 seed = 1
  complementary distribution
  response_levels = 3.6e+11 4.0e+11 4.4e+11
                   6.0e+04 6.5e+04 7.0e+04
                   3.5e+05 4.0e+05 4.5e+05
  compute gen_reliabilities
```

7.2.74.2.2.4 system

- **Keywords Area**
- **method**
- **local_reliability**
- **response_levels**
- **compute**
- **system**

Compute system reliability (series or parallel)

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>System Reliability Mode (Group 1)</td>
<td>series</td>
<td>Aggregate response statistics assuming a series system</td>
</tr>
</tbody>
</table>
Description

With the system probability/reliability option, statistics for specified response_levels are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

7.2.74.2.2.5  series

- Keywords Area
- method
- local_reliability
- response_levels
- compute
- system
- series

Aggregate response statistics assuming a series system

Specification

Alias: none

Argument(s): none

Description

See parent keyword system for description.

7.2.74.2.2.6  parallel

- Keywords Area
- method
- local_reliability
- response_levels
- compute
- system
- parallel

Aggregate response statistics assuming a parallel system
7.2.74.3 probability_levels

- Keywords Area
- method
- local_reliability
- probability_levels

Specify probability levels at which to estimate the corresponding response value

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_probability_levels</td>
<td></td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

**Description**

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

**Expected Output**

If probability_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

**Examples**

The Dakota input file below specifies a sampling method with probability levels of interest.
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

**PDF for response_fn_1:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.4221494996e+11</td>
<td>5.1384774972e-12</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>4.0634975300e+11</td>
<td>5.1454122311e-12</td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>5.4196114379e+11</td>
<td>2.4334239039e-12</td>
</tr>
</tbody>
</table>

**PDF for response_fn_2:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>5.6511827775e+04</td>
<td>1.9839945149e-05</td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>6.1603813790e+04</td>
<td>5.8916108390e-05</td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>7.8702465575e+04</td>
<td>2.9242071306e-05</td>
</tr>
</tbody>
</table>

**PDF for response_fn_3:**

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.6997214153e+05</td>
<td>5.302838523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.4111498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

**Level mappings for each response function:**

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
</tr>
</tbody>
</table>
7.2 method

4.0634975300e+11 3.3000000000e-01
5.4196143799e+11 0.0000000000e+00

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>7.8702457555e+04</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>1.0000000000e+00</td>
<td>1.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>3.8100966325e+05</td>
<td>2.0000000000e-01</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
<tr>
<td>4.4111498127e+05</td>
<td>0.0000000000e+00</td>
<td>0.0000000000e+00</td>
<td></td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.74.3.1 num_probability_levels

- Keywords Area
- method
- local_reliability
- probability_levels
- num_probability_levels

Specify which probability_levels correspond to which response

Specification

Alias: none
Argument(s): INTEGERLIST
Default: probability_levels evenly distributed among response functions

Description

See parent page

7.2.74.4 reliability_levels

- Keywords Area
- method
• local_reliability
• reliability_levels

Specify reliability levels at which the response values will be estimated

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_reliability_levels</td>
<td>Specify which reliability_levels correspond to which response</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF reliabilities by projecting out the prescribed number of sample standard deviations from the sample mean.

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.74.4.1 num_reliability_levels

• Keywords Area
• method
• local_reliability
• reliability_levels
• num_reliability_levels

Specify which reliability_levels correspond to which response

Specification

Alias: none
### Argument(s): INTEGERLIST

**Default:** reliability_levels evenly distributed among response functions

#### Description

See parent page

#### 7.2.74.5 gen_reliability_levels

- **Keywords Area**
- **method**
- **local_reliability**
- **gen_reliability_levels**

Specify generalized reliability levels at which to estimate the corresponding response value

#### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
<td></td>
</tr>
</tbody>
</table>

#### Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

#### Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).
7.2.74.5.1 num_gen_reliability_levels

- Keywords Area
- method
- local_reliability
- gen_reliability_levels
- num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** gen_reliability_levels evenly distributed among response functions

**Description**

See parent page

7.2.74.6 distribution

- Keywords Area
- method
- local_reliability
- distribution

Selection of cumulative or complementary cumulative functions

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** cumulative (CDF)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>cumulative</td>
<td>complementary</td>
<td>Computes statistics according to cumulative functions</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>
Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.

Default Behavior

If the distribution keyword is present, it must be accompanied by either cumulative or complementary. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.74.6.1 cumulative

- Keywords Area
- method
- local_reliability
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the cumulative keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.
Examples

method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative

7.2.74.2 complementary

  • Keywords Area
  • method
  • local_reliability
  • distribution
  • complementary

Computes statistics according to complementary cumulative functions.

Specification

Alias: none

Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the `complementary` keyword must be appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    distribution complementary

7.2.74.7 max_iterations

  • Keywords Area
  • method
  • local_reliability
  • max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods.
7.2.74 method

Topics
This keyword is related to the topics:
  • method_independent_controls

Specification
Alias: none
Argument(s): INTEGER
Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description
The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior
Default value is 100.

See Also
These keywords may also be of interest:
  • max_function_evaluations

7.2.74.8 convergence_tolerance

  • Keywords Area
  • method
  • local_reliability
  • convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics
This keyword is related to the topics:
  • method_independent_controls

Specification
Alias: none
Argument(s): REAL
Default: 1.e-4
Description

The `convergence_tolerance` specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration. Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.74.9 final_moments

- Keywords Area
- method
- local_reliability
- final_moments

Output moments of the specified type and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
Default: standard
Child Keywords:
7.2 method

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Objective Formulation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>none</td>
<td>standard</td>
<td>Output standardized moments and include them within the set of final statistics.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Output central moments and include them within the set of final statistics.</td>
</tr>
</tbody>
</table>

**Description**

When performing a nested study that may employ moment statistics on the inner loop, it can be desirable to control the type of these moments. The `final_moments` specification supports options of `none`, `standard` (default), or `central`, corresponding to omission of moments, standardized moments (mean, standard deviation, skewness, and excess kurtosis), or central moments (mean, variance, 3rd central, and 4th central).

The presence or omission of moment results in the final statistics influences the outer level mappings in the case of a nested study. For example, `final_moments none` can allow for a more compact specification of primary and/or secondary response mappings.

**Examples**

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments central
```

Overriding the default to `none` as follows:

```plaintext
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none
```

allows associated nested model mappings to be simplified from:

```plaintext
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 0. 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```

To a more compact version focused only on the response level mappings (two leading zeros per response function for moment mappings have been removed):

```plaintext
model
  nested
    primary_response_mapping = 0. 1. 0. 0. 1. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 0. 0. 0. 0. 0. 1.
```
model
  nested
    sub_method_pointer = 'UQ'
    primary_response_mapping = 1. 1. 0.
    secondary_response_mapping = 0. 0. 1.

7.2.74.9.1 none
  • Keywords Area
  • method
  • local_reliability
  • final_moments
  • none

Omit moments from the set of final statistics.

Specification

Alias: none
Argument(s): none

Description

The omission of moment results from the final statistics can allow for a more compact definition of primary and/or secondary response mappings within a nested model specification.

Examples

method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
    final_moments none

7.2.74.9.2 standard
  • Keywords Area
  • method
  • local_reliability
  • final_moments
  • standard

Output standardized moments and include them within the set of final statistics.

Specification

Alias: none
Argument(s): none
7.2 method

Description

Output standardized moments (mean, standard deviation, skewness, and excess kurtosis) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```

Output central moments and include them within the set of final statistics.

Specification

Alias: none

Argument(s): none

Description

Output central moments (mean, variance, 3rd central, and 4th central) and include the first two within the set of final statistics to be used at a higher level (e.g., optimization under uncertainty, mixed aleatory-epistemic UQ). This inclusion of moment results affects the primary and/or secondary response mappings in a nested model specification.

Examples

The following method specification overrides the default to print central moments and include them in the set of final statistics.

```
method,
  sampling
    samples = 50 seed = 1234
    response_levels = 3.6e+11 1.2e+05 3.5e+05
  final_moments central
```
7.2.74.10  model_pointer

- Keywords Area
- method
- local_reliability
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

**Examples**

```
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling, 
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                    0.1 0.2 0.6
                    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
```
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.2.75  global_reliability

• Keywords Area
• method
• global_reliability

Global reliability methods

Topics
This keyword is related to the topics:
• uncertainty_quantification
• reliability_methods

Specification
Alias: nond_global_reliability
Argument(s): none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>initial_samples</td>
<td>Initial number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Approximation (Group 1)</td>
<td>x_gaussian_process</td>
<td>Create GP surrogate in x-space</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>GP Implementation (Group 2)</td>
<td>u_gaussian_process</td>
<td>Create GP surrogate in u-space</td>
</tr>
<tr>
<td>Optional</td>
<td>GP Implementation</td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
<tr>
<td></td>
<td></td>
<td>dakota</td>
<td>Select the built in Gaussian Process surrogate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>experimental</td>
<td>Use the experimental Gaussian Process surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build_points_file</td>
<td></td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td></td>
<td>Output file for surrogate model value evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>use_derivatives</td>
<td></td>
<td>Use derivative data to construct surrogate models</td>
</tr>
<tr>
<td>Optional</td>
<td>seed</td>
<td></td>
<td>Seed of the random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>rng</td>
<td></td>
<td>Selection of a random number generator</td>
</tr>
<tr>
<td>Optional</td>
<td>response_levels</td>
<td></td>
<td>Values at which to estimate desired statistics for each response</td>
</tr>
</tbody>
</table>
### Description

These methods do not support forward/inverse mappings involving `reliability_levels`, since they never form a reliability index based on distance in u-space. Rather, they use a Gaussian process model to form an approximation to the limit state (based either in x-space via the `x_gaussian_process` specification or in u-space via the `u_gaussian_process` specification), followed by probability estimation based on multimodal adaptive importance sampling (see [10]) and [11]). These probability estimates may then be transformed into generalized reliability levels if desired. At this time, inverse reliability analysis (mapping probability or generalized reliability levels into response levels) is not implemented.

The Gaussian process model approximation to the limit state is formed over the aleatory uncertain variables by default, but may be extended to also capture the effect of design, epistemic uncertain, and state variables. If this is desired, one must use the appropriate controls to specify the active variables in the variables specification block. Refer to `variable_support` for additional information on supported variable types.

### See Also

These keywords may also be of interest:

- `adaptive_sampling`
- `gpais`
- `local_reliability`
- `sampling`
7.2.75.1  initial_samples

Initial number of samples for sampling-based methods

Specification

Alias: none

Argument(s): INTEGER

Default: \((d+1)(d+2)/2\)

Description

The initial_samples keyword is used to define the number of initial samples (i.e., randomly chosen sets of variable values) at which to execute a model. The initial samples may later be augmented in an iterative process.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least \(dim+1\) samples should be used, where "\(dim\)" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \((dim+1)(dim+2)/2\) samples are needed. For uncertainty quantification, we recommend at least \(10\times dim\) samples. For variance_based_decomp, we recommend hundreds to thousands of samples. Note that for variance_based_decomp, the number of simulations performed will be \(N\times(dim+2)\).

Examples

```
method sampling
  sample_type random
  initial_samples = 20
  refinement_samples = 5
```

7.2.75.2  x_gaussian_process

Create GP surrogate in x-space
Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: x_kriging
Argument(s): none

7.2.75.3 u_gaussian_process

- Keywords Area
- method
- global_reliability
- u_gaussian_process

Create GP surrogate in u-space

Topics

This keyword is related to the topics:

- reliability_methods

Specification

Alias: u_kriging
Argument(s): none

7.2.75.4 surfpack

- Keywords Area
- method
- global_reliability
- surfpack

Use the Surfpack version of Gaussian Process surrogates

Specification

Alias: none
Argument(s): none

Child Keywords:
### Description

This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. **Optimization methods**:
   - Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default, the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See `optimization_method`.
   - The total number of evaluations of the likelihood function can be controlled using the `max_trials` keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. **Trend Function**:
   - The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See `trend`.

3. **Correlation Lengths**:
   - Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See `correlation_lengths`.

4. **Ill-conditioning**
   - One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   - Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   - The `surfpack` model handles ill-conditioning internally by default, but behavior can be modified using `surfpack`.

5. **Gradient Enhanced Kriging (GEK)**
   - The `use_derivatives` keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   - See notes in the Theory section.

### Theory

**Gradient Enhanced Kriging**

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.
This issue was addressed by using a pivoted Cholesky factorization of Kriging’s correlation matrix (which is a small sub-matrix within GEK’s correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK’s correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging’s correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK’s correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK’s correlation matrix.

7.2.7.5.4.1 export_model

- Keywords Area
- method
- global_reliability
- surfpack
- export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>formats</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- text_archive: Plain-text, machine-readable archive for re-import or use with the surfpack executable
These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior

No export.

Expected Output

Output depends on selected format; see the formats specification.

Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern \{prefix\}.{response_descriptor}.{extension}. This filename_prefix keyword is used to supply the prefix portion of the pattern.

The response_descriptor portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns my_surrogate.{response_descriptor}.alg (for the algebraic files) and my_surrogate.{response_descriptor}.bsps (for the binary files).

```
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive
```

7.2.75.4.1.2 formats

- Keywords Area
- method
- global_reliability
- surfpack
- export_model
- formats

Formats for surrogate model export

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:
### Keywords Area

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</tr>
</thead>
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<td></td>
<td>Surrogate model</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>plain-text archive</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td></td>
<td>Surrogate model</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>binary archive file</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>format</td>
</tr>
</tbody>
</table>

#### Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted.

See `export_model` and the entries for the format selection keywords for further information.

7.2.75.4.1.3  text_archive

- Keywords Area
- method
- global_reliability
- surfpack
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format

#### Topics

This keyword is related to the topics:

- surrogate_models

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`.

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.
Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.75.4.1.4  binary_archive

- Keywords Area
- method
- global_reliability
- surfpack
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which 'bsps' stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.75.5  dakota

- Keywords Area
- method
- global_reliability
- dakota

Select the built in Gaussian Process surrogate
Specification

Alias: none
Argument(s): none

Description

A second version of GP surrogates was available in prior versions of Dakota. For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in[60].

Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a point_selection option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the point_selection option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

7.2.75.6 experimental

- Keywords Area
- method
- global_reliability
- experimental

Use the experimental Gaussian Process surrogate

Specification

Alias: none
Argument(s): none
Child Keywords:
### Description

The experimental GP surrogate is an emerging capability intended to supersede the `dakota` and `surfpack` GP implementations by unifying the best features of both. It also includes additional features such as the ability to export the model and then use it from Python.

### See Also

These keywords may also be of interest:

- `experimental_gaussian_process`

### 7.2.75.6.1 export_model

Exports surrogate model in user-specified format(s)

### Topics

This keyword is related to the topics:

- `surrogate_models`

### Specification

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>export_model</td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>options_file</td>
<td>Filename for a YAML file that specifies Gaussian process options</td>
</tr>
</tbody>
</table>
Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (`bin/surfpack`) or a user-developed tool. Export format is controlled using the `formats` specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- `text_archive` - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- `binary_archive` - Binary, machine-readable archive for re-import or use with the surfpack executable
- `algebraic_file` - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- `algebraic_console` - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process` `surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Funtions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only text_archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior

No export.

Expected Output

Output depends on selected format; see the `formats` specification.

Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.75.6.1.1 filename_prefix

- Keywords Area
- method
- global_reliability
- experimental
7.2 method

• export_model

• filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern {prefix}.{response_descriptor}.{extension}. This filename_prefix keyword is used to supply the prefix portion of the pattern.

The response_descriptor portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns my_surrogate.{response_descriptor}.alg (for the algebraic files) and my_surrogate.{response_descriptor}.bsps (for the binary files).

surrogate global gaussian_process surfpack
export_model
    filename_prefix = ’my_surrogate’
    formats
        algebraic_file
        binary_archive

7.2.75.6.1.2 formats

• Keywords Area

• method

• global_reliability

• experimental

• export_model

• formats

Formats for surrogate model export
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>text_archive</td>
<td></td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td></td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See export_model and the entries for the format selection keywords for further information.

7.2.75.6.1.3 text_archive

- Keywords Area
- method
- global_reliability
- experimental
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none
Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named \{prefix\}.{response_descriptor}.txt. Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.sps, in which ‘sps’ stands for Surfpack surrogate. See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.2.75.6.1.4 binary_archive

- Keywords Area
- method
- global_reliability
- experimental
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin. Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which ‘bsps’ stands for binary Surfpack surrogate. See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
7.2.75.6.2 options_file

- Keywords Area
- method
- global_reliability
- experimental
- options_file

Filename for a YAML file that specifies Gaussian process options

**Specification**

**Alias:** none  
**Argument(s):** STRING  
**Default:** no advanced options file

**Description**

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

**Examples**

```
# FILE: gp_options.yaml

GP Parameters:
  Sigma Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
  scaler name: standardization
  num restarts: 20
Nugget:
  fixed nugget: 0.0
  estimate nugget: true
  Bounds:
    lower bound: 3.17e-8
    upper bound: 1.0e-2
gp seed: 42
standardize response: false
Trend:
  estimate trend: true
  Options:
    max degree: 2
    reduced basis: false
    p-norm: 1.0
    scaler type: none
    regression solver type: SVD
    verbosity: 1
kernel type: squared exponential
Length-scale Bounds:
  lower bound: 1.0e-2
  upper bound: 1.0e2
  verbosity: 1
```

7.2.75.7 import_build_points_file

- Keywords Area
7.2 method

- method
- global_reliability
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

Alias: import_points_file

Argument(s): STRING

Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

method
polynomial_chaos
    expansion_order = 4
    import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'

7.2.75.7.1 custom_annotated

- Keywords Area
- method
- global_reliability
- import_build_points_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**
The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
   tabular_data
      tabular_data_file = 'dakota_summary.dat'
      custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id  x1   x2   obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  0.9   1.1   0.0002  0.26   0.76
2  0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3  0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...```

7.2.75.7.1.1 header

- Keywords Area
- method
- global_reliability
- import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom_annotated
7.2.75.7.1.2 eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no eval_id column

Description

See description of parent custom_annotated

7.2.75.7.1.3 interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent custom_annotated
7.2.75.7.2  annotated

- Keywords Area
- method
- global_reliability
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to `annotated` format, though `freeform` remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the `freeform` option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>@eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.75.7.3 freeform

- Keywords Area
- method
- global_reliability
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.
In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
 0.9  1.1 0.0002  0.26  0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.2.75.7.4 active_only

- Keywords Area
- method
- global_reliability
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.
7.2.75.8 export_approx_points_file

- Keywords Area
- method
- global_reliability
- export_approx_points_file

Output file for surrogate model value evaluations

**Specification**

**Alias:** export_points_file

**Argument(s):** STRING

**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

7.2.75.8.1 custom_annotated

- Keywords Area
- method
- global_reliability
- export_approx_points_file
- custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
### custom_annotated header eval_id

Enable header row in custom-annotated tabular file

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Default**: no header

**Description**

See description of parent `custom_annotated`

### eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

- **Alias**: none
- **Argument(s)**: none
- **Default**: no eval_id column
Description

See description of parent `custom_annotated`

7.2.75.8.1.3  `interface_id`

- Keywords Area
- method
- global_reliability
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no interface_id column

Description

See description of parent `custom_annotated`

7.2.75.8.2  `annotated`

- Keywords Area
- method
- global_reliability
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format
An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior
By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

• To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1  NO_ID  0.9  1.1  0.0002  0.26  0.76
2  NO_ID  0.90009  1.1  0.0001996404857  0.2601620081  0.759955
3  NO_ID  0.89991  1.1  0.0002003604863  0.2598380081  0.760045
...
```

7.2.75.8.3 freeform

• Keywords Area

• method

• global_reliability

• export_approx_points_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats
7.2 method

Specification

Alias: none

Argument(s): none

Default: annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
```

7.2.75.9 use_derivatives

- Keywords Area
- method
- global_reliability
- use_derivatives

Use derivative data to construct surrogate models
Specification

**Alias:** none

**Argument(s):** none

**Default:** use function values only

**Description**

The `use_derivatives` flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it’s use with Surfpack Gaussian process is not recommended.

7.2.75.10 seed

- **Keywords Area**
- **method**
- **global_reliability**
- **seed**

Seed of the random number generator

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** system-generated (non-repeatable)

**Description**

The random `seed` control provides a mechanism for making a stochastic method repeatable. That is, the use of the same random seed in identical studies will generate identical results.

**Default Behavior**

If not specified, the seed is randomly generated.

**Expected Output**

If `seed` is specified, a stochastic study will generate identical results when repeated using the same seed value. Otherwise, results are not guaranteed to be the same.

**Usage Tips**

If a stochastic study was run without `seed` specified, and the user later wishes to repeat the study using the same seed, the value of the seed used in the original study can be found in the output Dakota prints to the screen. That value can then be added to the Dakota input file.

**Examples**

```
method
  sampling
    sample_type lhs
```
samples = 10
seed = 15347

7.2.75.11  rng

- Keywords Area
- method
- global_reliability
- rng

Selection of a random number generator

Specification

Alias: none
Argument(s): none
Default: Mersenne twister (mt19937)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group RNG Algorithm (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>mt19937</td>
<td>Generates random numbers using the Mersenne twister</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rnum2</td>
<td>Generates pseudo-random numbers using the Pecos package</td>
</tr>
</tbody>
</table>

Description

The rng keyword is used to indicate a choice of random number generator.

Default Behavior

If specified, the rng keyword must be accompanied by either rnum2 (pseudo-random numbers) or mt19937 (random numbers generated by the Mersenne twister). Otherwise, mt19937, the Mersenne twister is used by default.

Usage Tips

The default is recommended, as the Mersenne twister is a higher quality random number generator.

Examples

method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
7.2.75.11.1 mt19937

- Keywords Area
- method
- global_reliability
- rng
- mt19937

Generates random numbers using the Mersenne twister

Specification

**Alias:** none

**Argument(s):** none

Description

The `mt19937` keyword directs Dakota to use the Mersenne twister to generate random numbers. Additional information can be found on wikipedia: [http://en.wikipedia.org/wiki/Mersenne_twister](http://en.wikipedia.org/wiki/Mersenne_twister).

Default Behavior

mt19937 is the default random number generator. To specify it explicitly in the Dakota input file, however, it must be specified in conjunction with the `rng` keyword.

Usage Tips

Use of the Mersenne twister random number generator (`mt19937`) is recommended.

Examples

```dakota
method
  sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng mt19937
```

7.2.75.11.2 rnum2

- Keywords Area
- method
- global_reliability
- rng
- rnum2

Generates pseudo-random numbers using the Pecos package

Specification

**Alias:** none

**Argument(s):** none
Description

The rnum2 keyword directs Dakota to use pseudo-random numbers generated by the Pecos package.

Default Behavior

rnum2 is not used by default. To change this behavior, it must be specified in conjunction with the rng keyword.

Usage Tips

Use of the Mersenne twister random number generator (mt19937) is recommended over rnum2.

Examples

```
method
sampling
    sample_type lhs
    samples = 10
    seed = 98765
    rng rnum2
```

7.2.75.12 response_levels

- Keywords Area
- method
- global_reliability
- response_levels

Values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF probabilities/reliabilities to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_response_levels</td>
<td>Number of values at which to estimate desired statistics for each response</td>
<td></td>
<td></td>
</tr>
<tr>
<td>compute</td>
<td>Selection of statistics to compute at each response</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Description

The response_levels specification provides the target response values for which to compute probabilities, reliabilities, or generalized reliabilities (forward mapping).
Default Behavior

If response_levels are not specified, no statistics will be computed. If they are, probabilities will be computed by default.

Expected Outputs

If response_levels are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the probability_levels and/or response_levels in the Dakota input file. If there are not very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Usage Tips

The num_response_levels is used to specify which arguments of the response_level correspond to which response.

Examples

For example, specifying a response_level of 52.3 followed with compute probabilities will result in the calculation of the probability that the response value is less than or equal to 52.3, given the uncertain distributions on the inputs.

For an example with multiple responses, the following specification

response_levels = 1. 2. .1 .2 .3 .4 10. 20. 30.
num_response_levels = 2 4 3

would assign the first two response levels (1., 2.) to response function 1, the next four response levels (.1, .2, .3, .4) to response function 2, and the final three response levels (10., 20., 30.) to response function 3. If the num_response_levels key were omitted from this example, then the response levels would be evenly distributed among the response functions (three levels each in this case).

The Dakota input file below specifies a sampling method with response levels of interest.

method, sampling,
samples = 100 seed = 1
complementary distribution
response_levels = 3.6e+11 4.0e+11 4.4e+11
6.0e+04 6.5e+04 7.0e+04
3.5e+05 4.0e+05 4.5e+05

variables,
  normal_uncertain = 2
  means = 248.89, 593.33
  std_deviations = 12.4, 29.7
  descriptors = 'TF1n' 'TF2n'
  uniform_uncertain = 2
  lower_bounds = 199.3, 474.63
  upper_bounds = 298.5, 712.
  descriptors = 'TF1u' 'TF2u'
  weibull_uncertain = 2
  alphas = 12., 30.
  betas = 250., 590.
  descriptors = 'TF1w' 'TF2w'
  histogram_bin_uncertain = 2
  num_pairs = 3 4
  abscissas = 5 8 10 .1 .2 .3 .4
  counts = 17 21 0 12 24 12 0
  descriptors = 'TF1h' 'TF2h'
  histogram_point_uncertain
  real = 1
  num_pairs = 2
  abscissas = 3 4
Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values specified in the input file. The probability levels corresponding to those response values are shown in the CCDF.

Probability Density Function (PDF) histograms for each response function:

PDF for response_fn_1:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>3.6000000000e+11</td>
<td>5.361733194e-12</td>
</tr>
<tr>
<td>3.6000000000e+11</td>
<td>4.0000000000e+11</td>
<td>4.2500000000e-12</td>
</tr>
<tr>
<td>4.0000000000e+11</td>
<td>4.4000000000e+11</td>
<td>3.7500000000e-12</td>
</tr>
<tr>
<td>4.4000000000e+11</td>
<td>5.4196114379e+11</td>
<td>2.2557612778e-12</td>
</tr>
</tbody>
</table>

PDF for response_fn_2:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>6.0000000000e+04</td>
<td>2.8742313192e-05</td>
</tr>
<tr>
<td>6.0000000000e+04</td>
<td>6.5000000000e+04</td>
<td>6.4000000000e-05</td>
</tr>
<tr>
<td>6.5000000000e+04</td>
<td>7.0000000000e+04</td>
<td>4.0000000000e-05</td>
</tr>
<tr>
<td>7.0000000000e+04</td>
<td>7.8702465755e+04</td>
<td>1.0341896485e-05</td>
</tr>
</tbody>
</table>

PDF for response_fn_3:

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.3796737090e+05</td>
<td>3.5000000000e+05</td>
<td>4.2844660868e-06</td>
</tr>
<tr>
<td>3.5000000000e+05</td>
<td>4.0000000000e+05</td>
<td>8.6000000000e-06</td>
</tr>
<tr>
<td>4.0000000000e+05</td>
<td>4.5000000000e+05</td>
<td>1.8000000000e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

Complementary Cumulative Distribution Function (CCDF) for response_fn_1:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.6000000000e+11</td>
<td>5.0000000000e-01</td>
<td>3.8000000000e-01</td>
<td>2.3000000000e-01</td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_2:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>6.0000000000e+04</td>
<td>6.1000000000e-01</td>
<td>2.9000000000e-01</td>
<td>9.0000000000e-02</td>
</tr>
</tbody>
</table>

Complementary Cumulative Distribution Function (CCDF) for response_fn_3:

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.5000000000e+05</td>
<td>9.0000000000e-02</td>
<td>9.0000000000e-02</td>
<td>9.0000000000e-02</td>
</tr>
</tbody>
</table>

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

A forward mapping involves computing the belief and plausibility probability level for a specified response level.
7.2.75.12.1  num_response_levels

- Keywords Area
- method
- global_reliability
- response_levels
- num_response_levels

Number of values at which to estimate desired statistics for each response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: response_levels evenly distributed among response functions

Description

The `num_response_levels` keyword allows the user to specify the number of response values, for each response, at which estimated statistics are of interest. Statistics that can be computed are probabilities and reliabilities, both according to either a cumulative distribution function or a complementary cumulative distribution function.

Default Behavior

If `num_response_levels` is not specified, the response_levels will be evenly distributed among the responses.

Expected Outputs

The specific output will be determined by the type of statistics that are specified. In a general sense, the output will be a list of response level-statistic pairs that show the estimated value of the desired statistic for each response level specified.

Examples

```yaml
method
  sampling
    samples = 100
    seed = 34785
    num_response_levels = 1 1 1
    response_levels = 0.5 0.5 0.5
```

7.2.75.12.2  compute

- Keywords Area
- method
- global_reliability
- response_levels
- compute

Selection of statistics to compute at each response level
7.2 method

Specification

**Alias:** none

**Argument(s):** none

**Default:** probabilities

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Statistics to Compute (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>probabilities</td>
<td>Computes probabilities associated with response levels</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>gen_reliabilities</td>
<td>Computes generalized reliabilities associated with response levels</td>
</tr>
<tr>
<td></td>
<td></td>
<td>system</td>
<td>Compute system reliability (series or parallel)</td>
</tr>
</tbody>
</table>

**Description**

The `compute` keyword is used to select which forward statistical mapping is calculated at each response level.

**Default Behavior**

If `response_levels` is not specified, no statistics are computed. If `response_levels` is specified but `compute` is not, probabilities will be computed by default. If both `response_levels` and `compute` are specified, then one of the following must be specified: `probabilities`, `reliabilities`, or `gen_reliabilities`.

**Expected Output**

The type of statistics specified by `compute` will be reported for each response level.

**Usage Tips**

CDF/CCDF probabilities are calculated for specified response levels using a simple binning approach.

CDF/CCDF reliabilities are calculated for specified response levels by computing the number of sample standard deviations separating the sample mean from the response level.

**Examples**

```plaintext
method
    sampling
        sample_type random
        samples = 100 seed = 1
        complementary distribution
        response_levels = 3.6e+11 4.0e+11 4.4e+11 6.0e+04 6.5e+04 7.0e+04 3.5e+05 4.0e+05 4.5e+05
    compute reliabilities
```
7.2.75.12.2.1 probabilities

- Keywords Area
- method
- global_reliability
- response_levels
- compute
- probabilities

Computes probabilities associated with response levels

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The `probabilities` keyword directs Dakota to compute the probability that the model response will be below (cumulative) or above (complementary cumulative) a specified response value. This is done for every response level designated for each response.

**Default Behavior**

If `response_levels` is specified, the probabilities are computed by default. To explicitly specify it in the Dakota input file, though, the `probabilities` keyword should be specified in conjunction with the `compute` keyword.

**Expected Outputs**

The Dakota output is a set of response level-probability pairs that give the probability that the model response will be below or above the corresponding response level, depending on the distribution defined.

**Examples**

```
method sampling
    sample_type random
    samples = 100 seed = 1
    complementary distribution
    response_levels = 3.6e+11 4.0e+11 4.4e+11
                      6.0e+04 6.5e+04 7.0e+04
                      3.5e+05 4.0e+05 4.5e+05
    compute probabilities
```

7.2.75.12.2.2 gen_reliabilities

- Keywords Area
- method
- global_reliability
- response_levels
- compute
- gen_reliabilities

Computes generalized reliabilities associated with response levels
7.2 method 3225

 Specification

 Alias: none
 Argument(s): none

 Description

 The gen_reliabilities keyword directs Dakota to compute generalized reliabilities according to the specified distribution for a specified response value. This is done for every response level designated for each response.

 Default Behavior

 If response_levels is specified, the generalized reliabilities are not computed by default. To change this behavior, the gen_reliabilities keyword should be specified in conjunction with the compute keyword.

 Expected Outputs

 The Dakota output is a set of response level-generalized reliability pairs according to the distribution defined.

 Examples

 method sampling
     sample_type random
     samples = 100 seed = 1
     complementary_distribution
     response_levels = 3.6e+11 4.0e+11 4.4e+11
                       6.0e+04 6.5e+04 7.0e+04
                       3.5e+05 4.0e+05 4.5e+05
     compute gen_reliabilities

 7.2.75.12.2.3 system

  • Keywords Area

  • method

  • global_reliability

  • response_levels

  • compute

  • system

 Compute system reliability (series or parallel)

 Specification

 Alias: none
 Argument(s): none
 Child Keywords:
### Description

With the system probability/reliability option, statistics for specified `response_levels` are calculated and reported assuming the response functions combine either in series or parallel to produce a total system response.

For a series system, the system fails when any one component (response) fails. The probability of failure is the complement of the product of the individual response success probabilities.

For a parallel system, the system fails only when all components (responses) fail. The probability of failure is the product of the individual response failure probabilities.

#### 7.2.75.12.2.4 series

- **Keywords Area**
- `method`
- `global_reliability`
- `response_levels`
- `compute`
- `system`
- `series`

Aggregate response statistics assuming a series system

#### Specification

**Alias:** none

**Argument(s):** none

#### Description

See parent keyword `system` for description.

#### 7.2.75.12.2.5 parallel

- **Keywords Area**
- `method`
- `global_reliability`
- `response_levels`
7.2 method

- compute
- system
- parallel

Aggregate response statistics assuming a parallel system

Specification

**Alias:** none  
**Argument(s):** none

Description

See parent keyword `system` for description.

7.2.75.13 probability_levels

- Keywords Area
- method
- global_reliability
- probability_levels

Specify probability levels at which to estimate the corresponding response value

Specification

**Alias:** none  
**Argument(s):** REALLIST

**Default:** No CDF/CCDF response levels to compute

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_probability_levels</td>
<td>Specify which probability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified CDF/CCDF probabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Expected Output

If `probability_levels` are specified, Dakota will create two tables in the standard output: a Probability Density function (PDF) histogram and a Cumulative Distribution Function (CDF) table. The PDF histogram has the lower and upper endpoints of each bin and the corresponding density of that bin. Note that the PDF histogram has bins defined by the `probability_levels` and/or `response_levels` in the Dakota input file. If there are not
very many levels, the histogram will be coarse. Dakota does not do anything to optimize the bin size or spacing. The CDF table has the list of response levels and the corresponding probability that the response value is less than or equal to each response level threshold.

Examples

The Dakota input file below specifies a sampling method with probability levels of interest.

```
method,
    sampling,
    samples = 100 seed = 1
    complementary_distribution
    probability_levels = 1. .66 .33 0.
    1. .8 .5 0.
    1. .3 .2 0.
variables,
    normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
    uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
    weibull_uncertain = 2
    alphas = 12., 30.
    betas = 250., 590.
    descriptors = 'TF1w' 'TF2w'
    histogram_bin_uncertain = 2
    num_pairs = 3 4
    abscissas = 5, 8, 10 .1 .2 .3 .4
    counts = 17, 21 0 12 24 12 0
    descriptors = 'TF1h' 'TF2h'
    histogram_point_uncertain
    real = 1
    num_pairs = 2
    abscissas = 3 4
    counts = 1 1
    descriptors = 'TF3h'
interface,
    system async evaluation_concurrency = 5
    analysis_driver = 'text_book'
responses,
    response_functions = 3
    no_gradients
    no_hessians
```

Given the above Dakota input file, the following excerpt from the output shows the PDF and CCDF generated. Note that the bounds on the bins of the PDF are the response values that correspond the probability levels specified in the input file. Those response values are also shown in the CCDF.

<p>| Probability Density Function (PDF) histograms for each response function: |
|-------------------------------------------------|----------------|----------------|</p>
<table>
<thead>
<tr>
<th>PDF for response_fn_1:</th>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11 3.4221494996e+11 5.1384774972e-12</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11 4.0634975300e+11 2.4334239039e-12</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<p>| Probability Density Function (PDF) histograms for each response function: |
|-------------------------------------------------|----------------|----------------|</p>
<table>
<thead>
<tr>
<th>PDF for response_fn_2:</th>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04 5.6511827775e+04 1.9839945149e-05</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04 6.1603813790e+04 5.8916108390e-05</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04 7.8702465755e+04 2.9242071306e-05</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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7.2 method

<table>
<thead>
<tr>
<th>Bin Lower</th>
<th>Bin Upper</th>
<th>Density Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>3.6997214153e+05</td>
<td>5.3028386523e-06</td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.8100966235e+05</td>
<td>9.060055634e-06</td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>4.411498127e+05</td>
<td>3.3274925348e-06</td>
</tr>
</tbody>
</table>

Level mappings for each response function:

**Complementary Cumulative Distribution Function (CCDF) for response_fn_1:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.7604749078e+11</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.4221494996e+11</td>
<td>6.6000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.0634975300e+11</td>
<td>3.3000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.4196114379e+11</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_2:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.6431154744e+04</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5.6511827775e+04</td>
<td>8.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6.1603813790e+04</td>
<td>5.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.8702465755e+04</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Complementary Cumulative Distribution Function (CCDF) for response_fn_3:**

<table>
<thead>
<tr>
<th>Response Level</th>
<th>Probability Level</th>
<th>Reliability Index</th>
<th>General Rel Index</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.379673709e+05</td>
<td>1.0000000000e+00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.6997214153e+05</td>
<td>3.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3.8100966235e+05</td>
<td>2.0000000000e-01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4.411498127e+05</td>
<td>0.0000000000e+00</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Theory**

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.75.13.1 num_probability_levels

- **Keywords Area**
- **method**
- **global_reliability**
- **probability_levels**
- **num_probability_levels**

Specify which probability_levels correspond to which response

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** probability_levels evenly distributed among response functions
Description

See parent page

7.2.75.14  gen_reliability_levels

• Keywords Area
• method
• global_reliability
• gen_reliability_levels

Specify generalized reliability levels at which to estimate the corresponding response value

Specification

Alias: none

Argument(s): REALLIST

Default: No CDF/CCDF response levels to compute

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword Description</th>
<th>Dakota Keyword</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>num_gen_reliability_levels</td>
<td>Specify which gen_reliability_levels correspond to which response</td>
</tr>
</tbody>
</table>

Description

Response levels are calculated for specified generalized reliabilities by indexing into a sorted samples array (the response levels computed are not interpolated and will correspond to one of the sampled values).

Theory

Sets of response-probability pairs computed with the forward/inverse mappings define either a cumulative distribution function (CDF) or a complementary cumulative distribution function (CCDF) for each response function.

In the case of evidence-based epistemic methods, this is generalized to define either cumulative belief and plausibility functions (CBF and CPF) or complementary cumulative belief and plausibility functions (CCBF and CCPF) for each response function.

An inverse mapping involves computing the belief and plausibility response level for either a specified probability level or a specified generalized reliability level (two results for each level mapping in the evidence-based epistemic case, instead of the one result for each level mapping in the aleatory case).

7.2.75.14.1  num_gen_reliability_levels

• Keywords Area
• method
• global_reliability
• gen_reliability_levels
• num_gen_reliability_levels

Specify which gen_reliability_levels correspond to which response

Specification

Alias: none

Argument(s): INTEGERLIST

Default: gen_reliability_levels evenly distributed among response functions

Description

See parent page

7.2.75.15 distribution

• Keywords Area
• method
• global_reliability
• distribution

Selection of cumulative or complementary cumulative functions

Specification

Alias: none

Argument(s): none

Default: cumulative (CDF)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Distribution Type (CDF/CCDF) (Group 1)</th>
<th>Dakota Keyword cumulative</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>complementary</td>
<td>Computes statistics according to complementary cumulative functions</td>
</tr>
</tbody>
</table>

Description

The distribution keyword allows the user to select between a cumulative distribution/belief/plausibility function and a complementary cumulative distribution/belief/plausibility function. This choice affects how probabilities and reliability indices are reported.
Default Behavior

If the `distribution` keyword is present, it must be accompanied by either `cumulative` or `complementary`. Otherwise, a cumulative distribution will be used by default.

Expected Outputs

Output will be a set of model response-probability pairs determined according to the choice of distribution. The choice of distribution also defines the sign of the reliability or generalized reliability indices.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```

7.2.75.15.1 cumulative

- Keywords Area
- method
- global_reliability
- distribution
- cumulative

Computes statistics according to cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a cumulative distribution/belief/plausibility function.

Default Behavior

By default, a cumulative distribution/belief/plausibility function will be used. To explicitly specify it in the Dakota input file, however, the `cumulative` keyword must appear in conjunction with the `distribution` keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls below given response thresholds.

Examples

```
method
  sampling
    sample_type lhs
    samples = 10
    distribution cumulative
```
7.2.75.15.2 complementary

- Keywords Area
- method
- global_reliability
- distribution
- complementary

Computes statistics according to complementary cumulative functions

Specification

Alias: none
Argument(s): none

Description

Statistics on model responses will be computed according to a complementary cumulative distribution/belief/plausibility function.

Default Behavior

By default, a complementary cumulative distribution/belief/plausibility function will not be used. To change that behavior, the complementary keyword must be appear in conjunction with the distribution keyword.

Expected Outputs

Output will be a set of model response-probability pairs determined according to a complementary cumulative distribution/belief/plausibility function. The probabilities reported are the probabilities that the model response falls above given response thresholds.

Examples

```
method
sampling
  sample_type lhs
  samples = 10
  distribution complementary
```

7.2.75.16 max_iterations

- Keywords Area
- method
- global_reliability
- max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:
- method_independent_controls
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25+n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed $\text{max\_iterations}$ iterations. In cases where there are multiple nested iterations to be controlled, $\text{max\_iterations}$ generally controls the outer-most context.

**Default Behavior**

Default value is 100.

**See Also**

These keywords may also be of interest:

- max\_function\_evaluations

7.2.75.17 convergence_tolerance

- Keywords Area
- method
- global\_reliability
- convergence tolerance

Stopping criterion based on objective function or statistics convergence

**Topics**

This keyword is related to the topics:

- method\_independent\_controls

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** 1.e-4

**Description**

The $\text{convergence\_tolerance}$ specification provides a real value for controlling the termination of iteration.
For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.

Therefore, permissible values are between 0 and 1, non-inclusive.

Behavior Varies by Package/Library

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.75.18 model_pointer

- Keywords Area
- method
- global_reliability
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.
Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                      0.1 0.2 0.6
                      0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians
```

7.2.76 fsu_quasi_mc

- Keywords Area
- method
- fsu_quasi_mc

Design of Computer Experiments - Quasi-Monte Carlo sampling
7.2 method

Topics

This keyword is related to the topics:

- package_fsudace
- design_and_analysis_of_computer_experiments

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Type</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Sequence Type (Group 1)</td>
<td>halton</td>
<td>Generate samples from a Halton sequence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>hammersley</td>
<td>Use Hammersley sequences</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>latinize</td>
<td>Adjust samples to improve the discrepancy of the marginal distributions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>quality_metrics</td>
<td>Calculate metrics to assess the quality of quasi-Monte Carlo samples</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>variance_based_decomp</td>
<td>Activates global sensitivity analysis based on decomposition of response variance into contributions from variables</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>samples</td>
<td>Number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>fixed_sequence</td>
<td>Reuse the same sequence and samples for multiple sampling sets</td>
</tr>
</tbody>
</table>
Quasi-Monte Carlo methods produce low discrepancy sequences, especially if one is interested in the uniformity of projections of the point sets onto lower dimensional faces of the hypercube (usually 1-D: how well do the marginal distributions approximate a uniform?)

This method generates sets of uniform random variables on the interval [0,1]. If the user specifies lower and upper bounds for a variable, the [0,1] samples are mapped to the [lower, upper] interval.

The user must first choose the sequence type:

- halton
- hammersley

Then three keywords are used to define the sequence and how it is sampled:

- prime_base
- sequence_start
- sequence_leap

Each of these has defaults, so specification is optional.

Theory

The quasi-Monte Carlo sequences of Halton and Hammersley are deterministic sequences determined by a set of prime bases. Generally, we recommend that the user leave the default setting for the bases, which are the lowest primes. Thus, if one wants to generate a sample set for 3 random variables, the default bases used are 2, 3, and 5 in the Halton sequence. To give an example of how these sequences look, the Halton sequence in base 2 starts with points 0.5, 0.25, 0.75, 0.125, 0.625, etc. The first few points in a Halton base 3 sequence are 0.33333, 0.66667, 0.11111, 0.44444, 0.77777, etc. Notice that the Halton sequence tends to alternate back and forth, generating a point closer to zero then a point closer to one. An individual sequence is based on a radix inverse function defined on a prime base. The prime base determines how quickly the [0,1] interval is filled in. Generally, the lowest primes are recommended.

The Hammersley sequence is the same as the Halton sequence, except the values for the first random variable are equal to 1/N, where N is the number of samples. Thus, if one wants to generate a sample set of 100 samples for 3 samples...
random variables, the first random variable has values 1/100, 2/100, 3/100, etc. and the second and third variables are generated according to a Halton sequence with bases 2 and 3, respectively.

For more information about these sequences, see [44], [45], and [1].

See Also

These keywords may also be of interest:

- dace
- fsu_cvt
- psuade_moat

7.2.76.1 halton

- Keywords Area
- method
- fsu_quasi_mc
- halton

Generate samples from a Halton sequence

Topics

This keyword is related to the topics:

- package_fsudace

Specification

Alias: none
Argument(s): none

Description

The quasi-Monte Carlo sequences of Halton are deterministic sequences determined by a set of prime bases. These sequences generate random numbers with the goal of filling a unit hypercube uniformly.

Generally, we recommend that the user leave the default setting for the bases, which are the lowest primes. Thus, if one wants to generate a sample set for 3 random variables, the default bases used are 2, 3, and 5 in the Halton sequence. To give an example of how these sequences look, the Halton sequence in base 2 starts with points 0.5, 0.25, 0.75, 0.125, 0.625, etc. The first few points in a Halton base 3 sequence are 0.33333, 0.66667, 0.11111, 0.44444, 0.77777, etc. Notice that the Halton sequence tends to alternate back and forth, generating a point closer to zero then a point closer to one. An individual sequence is based on a radix inverse function defined on a prime base. The prime base determines how quickly the [0,1] interval is filled in.

Theory

For more information about these sequences, see [44], [45], and [1].
7.2.76.2 hammersley

- Keywords Area
- method
- fsu_quasi_mc
- hammersley

Use Hammersley sequences

Topics

This keyword is related to the topics:

- package_fsudace
- design_and_analysis_of_computer_experiments

Specification

Alias: none

Argument(s): none

Description

The Hammersley sequence is the same as the Halton sequence, except the values for the first random variable are equal to 1/N, where N is the number of samples. Thus, if one wants to generate a sample set of 100 samples for 3 random variables, the first random variable has values 1/100, 2/100, 3/100, etc. and the second and third variables are generated according to a Halton sequence with bases 2 and 3, respectively.

See Also

These keywords may also be of interest:

- fsu_quasi_mc

7.2.76.3 latinize

- Keywords Area
- method
- fsu_quasi_mc
- latinize

Adjust samples to improve the discrepancy of the marginal distributions

Specification

Alias: none

Argument(s): none

Default: No latinization
Description

The `latinize` control takes the samples and "latinizes" them, meaning that each original sample is moved so that it falls into one strata or bin in each dimension as in Latin Hypercube sampling. The default setting is NOT to latinize. However, one may be interested in doing this in situations where one wants better discrepancy of the 1-dimensional projections (the marginal distributions).

7.2.76.4 quality_metrics

- Keywords Area
- method
- fsu_quasi_mc
- quality_metrics

Calculate metrics to assess the quality of quasi-Monte Carlo samples

Topics

This keyword is related to the topics:

- package_fsudace

Specification

Alias: none
Argument(s): none
Default: No quality_metrics

Description

`quality_metrics` calculates four quality metrics relating to the volumetric spacing of the samples. The four quality metrics measure different aspects relating to the uniformity of point samples in hypercubes. Desirable properties of such point samples are:

- are the points equally spaced
- do the points cover the region
- and are they isotropically distributed
- with no directional bias in the spacing

The four quality metrics we report are:

- h: the point distribution norm, which is a measure of uniformity of the point distribution
- chi: a regularity measure, and provides a measure of local uniformity of a set of points
- tau: the second moment trace measure
- d: the second moment determinant measure

All of these values are scaled so that smaller is better (the smaller the metric, the better the uniformity of the point distribution).
Examples

Complete explanation of these measures can be found in [39].

7.2.76.5 variance_based_decomp

- Keywords Area
- method
- fsu_quasi_mc
- variance_based_decomp

Activates global sensitivity analysis based on decomposition of response variance into contributions from variables

Specification

Alias: none

Argument(s): none

Default: no variance-based decomposition

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>drop_tolerance</td>
<td>Suppresses output of sensitivity indices with values lower than this tolerance</td>
</tr>
</tbody>
</table>

Description

Dakota can calculate sensitivity indices through variance based decomposition using the keyword variance-based_decomp. These indicate how important the uncertainty in each input variable is in contributing to the output variance.

Default Behavior

Because of the computational cost, variance_based_decomp is turned off as a default.

If the user specified a number of samples, N, and a number of nondeterministic variables, M, variance-based decomposition requires the evaluation of $N \times (M + 2)$ samples. Note that specifying this keyword will increase the number of function evaluations above the number requested with the samples keyword since replicated sets of sample values are evaluated.

Expected Outputs

When variance_based_decomp is specified, sensitivity indices for main effects and total effects will be reported. Main effects (roughly) represent the percent contribution of each individual variable to the variance in the model response. Total effects represent the percent contribution of each individual variable in combination with all other variables to the variance in the model response.

Usage Tips

To obtain sensitivity indices that are reasonably accurate, we recommend that N, the number of samples, be at least one hundred and preferably several hundred or thousands.
Examples

```plaintext
method,
sampling
    sample_type lhs
    samples = 100
    variance_based_decomp
```

Theory

In this context, we take sensitivity analysis to be global, not local as when calculating derivatives of output variables with respect to input variables. Our definition is similar to that of \[76\]: “The study of how uncertainty in the output of a model can be apportioned to different sources of uncertainty in the model input.”

Variance based decomposition is a way of using sets of samples to understand how the variance of the output behaves, with respect to each input variable. A larger value of the sensitivity index, $S_i$, means that the uncertainty in the input variable $i$ has a larger effect on the variance of the output. More details on the calculations and interpretation of the sensitivity indices can be found in \[76\] and \[90\].

7.2.76.5.1 drop_tolerance

- Keywords Area
- method
- fsu_quasi_mc
- variance_based_decomp
- drop_tolerance

Suppresses output of sensitivity indices with values lower than this tolerance

Specification

**Alias:** none

**Argument(s):** REAL

**Default:** All VBD indices displayed

Description

The `drop_tolerance` keyword allows the user to specify a value below which sensitivity indices generated by `variance_based_decomp` are not displayed.

**Default Behavior**

By default, all sensitivity indices generated by `variance_based_decomp` are displayed.

**Usage Tips**

For `polynomial_chaos`, which outputs main, interaction, and total effects by default, the `univariate_effects` may be a more appropriate option. It allows suppression of the interaction effects since the output volume of these results can be prohibitive for high dimensional problems. Similar to suppression of these interactions is the covariance control, which can be selected to be diagonal_covariance or full_covariance, with the former supporting suppression of the off-diagonal covariance terms (to save compute and memory resources and reduce output volume).
Examples

```
method,
sampling
  sample_type lhs
  samples = 100
  variance_based_decomp
  drop_tolerance = 0.001
```

7.2.76.6 samples

- Keywords Area
- method
- fsu_quasi_mc
- samples

Number of samples for sampling-based methods

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 0

Description

The `samples` keyword is used to define the number of samples (i.e., randomly chosen sets of variable values) at which to execute a model.

**Default Behavior**

By default, Dakota will use the minimum number of samples required by the chosen method.

**Usage Tips**

To obtain linear sensitivities or to construct a linear response surface, at least dim+1 samples should be used, where "dim" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \((\text{dim}+1)(\text{dim}+2)/2\) samples are needed. For uncertainty quantification, we recommend at least 10\(^*\)dim samples. For `variance_based_decomp`, we recommend hundreds to thousands of samples. Note that for `variance_based_decomp`, the number of simulations performed will be N\(^*\)(dim+2).

Examples

```
method
  sampling
    sample_type lhs
    samples = 20
```

7.2.76.7 fixed_sequence

- Keywords Area
- method
- fsu_quasi_mc
7.2 method

- fixed_sequence

Reuse the same sequence and samples for multiple sampling sets

Specification

**Alias:** none

**Argument(s):** none

**Default:** sequence not fixed: sampling patterns are variable among multiple QMC runs

**Description**

The `fixed_sequence` control is similar to `fixed_seed` for other sampling methods. If `fixed_sequence` is specified, the user will get the same sequence (meaning the same set of samples) for subsequent calls of the QMC sampling method (for example, this might be used in a surrogate based optimization method or a parameter study where one wants to fix the uncertain variables).

7.2.76.8 sequence_start

- Keywords Area
- method
- fsu_quasi_mc
- sequence_start

Choose where to start sampling the sequence

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Default:** Vector of zeroes

**Description**

`sequence_start` determines where in the sequence the samples will start.

The default `sequence_start` is a vector with 0 for each variable, specifying that each sequence start with the first term.

**Examples**

For example, for the Halton sequence in base 2, if the user specifies `sequence_start = 2`, the sequence would not include 0.5 and 0.25, but instead would start at 0.75.

7.2.76.9 sequence_leap

- Keywords Area
- method
Specify how often the sequence is sampled

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** Vector of ones

**Description**

`sequence_leap` controls the "leaping" of terms in the sequence. The default is 1 for each variable, meaning that each term in the sequence be returned.

**Examples**

If the user specifies a `sequence_leap` of 2 for a variable, the points returned would be every other term from the QMC sequence.

**Theory**

The advantage to using a leap value greater than one is mainly for high-dimensional sets of random deviates. In this case, setting a leap value to the next prime number larger than the largest prime base can help maintain uniformity when generating sample sets for high dimensions. For more information about the efficacy of leaped Halton sequences, see[74].

7.2.76.10 prime_base

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST  
**Default:** Vector of the first s primes for s-dimensions in Halton, First (s-1) primes for Hammersley
7.2 method

Description

It is recommended that the user not specify this and use the default values.

• For the Halton sequence, the default bases are primes in increasing order, starting with 2, 3, 5, etc.

• For the Hammersley sequence, the user specifies (s-1) primes if one is generating an s-dimensional set of random variables.

7.2.76.11 max_iterations

• Keywords Area

• method

• fsu_quasi_mc

• max_iterations

Number of iterations allowed for optimizers and adaptive UQ methods

Topics

This keyword is related to the topics:

• method_independent_controls

Specification

Alias: none

Argument(s): INTEGER

Default: 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

Default Behavior

Default value is 100.

See Also

These keywords may also be of interest:

• max_function_evaluations
7.2.76.12  model_pointer

- Keywords Area
- method
- fsu_quasi_mc
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

evironment
    tabular_data
    method_pointer = 'UQ'

method
    id_method = 'UQ'
    model_pointer = 'SURR'
    sampling,
        samples = 10
        seed = 98765 rng rnum2
        response_levels = 0.1 0.2 0.6
            0.1 0.2 0.6
                0.1 0.2 0.6
        sample_type lhs
        distribution cumulative

model
    id_model = 'SURR'
7.2 method

surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.77 vector_parameter_study

• Keywords Area

• method

• vector_parameter_study

Samples variables along a user-defined vector

Topics

This keyword is related to the topics:

• parameter_studies

Specification

Alias: none
Argument(s): none
Child Keywords:
Description

Dakota’s vector parameter study computes response data sets at selected intervals along a vector in parameter space. It is often used for single-coordinate parameter studies (to study the effect of a single variable on a response set), but it can be used more generally for multiple coordinate vector studies (to investigate the response variations along some n-dimensional vector such as an optimizer search direction).

Default Behavior

By default, the vector parameter study operates over all types of variables.

Expected Outputs

A vector parameter study produces a set of responses for each parameter set that is generated.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Parameter Sets

Usage Tips

Group 1 is used to define the vector along which the parameters are varied. Both cases also rely on the variables specification of an initial value, through:

- the initial_point keyword
- the initial_state keyword
- relying on the default initial value, based on the rest of the variables specification

From the initial value, the vector can be defined using one of the two keyword choices. Once the initial vector is defined, the samples are then fully specified by num_steps.

Examples

The following example is a good comparison to the examples on multidim_parameter_study and centered_parameter_study.
# tested on Dakota 6.0 on 140501

environment
tabular_data
tabular_data_file = 'rosen_vector.dat'

method
vector_parameter_study
    num_steps = 10
    final_point = 2.0 2.0

model
    single

variables
    continuous_design = 2
    initial_point = -2.0 -2.0
descriptors = 'x1' 'x2'

interface
    analysis_driver = 'rosenbrock'
    fork

responses
    response_functions = 1
    no_gradients
    no_hessians

See Also

These keywords may also be of interest:

• centered_parameter_study
• multidim_parameter_study
• list_parameter_study

7.2.77.1 final_point

• Keywords Area
• method
• vector_parameter_study
• final_point

Final variable values defining vector in vector parameter study

Specification

Alias: none
Argument(s): REALLIST

Description

The final_point keyword is used to define the final values for each variable on the vector to be used in the vector parameter study. The vector’s direction and magnitude are determined by the initial value from the variables specification, and the final_point.

Default Behavior
The user is required to specify either final_point or step_vector. There is no default definition for the vector.

Usage Tips
The actual points are determined based on this vector and the number of points chosen is given in num_points.

Examples

```
method
  vector_parameter_study
  num_steps = 10
  final_point = 2.0 2.0
```

7.2.77.2 step_vector

- Keywords Area
- method
- vector_parameter_study
- step_vector

Size of step for each variable

Specification

Alias: none

Argument(s): REALLIST

Description

The step_vector keyword specifies how much each variable will be incremented in a single step. step_vector works in conjunction with num_steps, which determines the number of steps taken during the vector_parameter_study. If instead of step_vector, final_point is specified with num_steps, Dakota will infer the step sizes.

Entries in the step_vector are the actual amounts by which continuous and range variables are incremented. For set variables, step_vector entries are interpreted as indexes into the underlying set.

Default Behavior
The user is required to specify either final_point or step_vector. There is no default definition for the vector.

Examples

```
variables
  continuous_design 1
    initial_point 1.0
    descriptors 'x1'
  discrete_design_set
    string 1
      elements 'bar' 'baz' 'foo' 'fuzz'
    initial_point 'bar'
    descriptors 's1'
method
```
vector_parameter_study
  num_steps = 3
  # Add 2.0 to x1 and increment s1 by 1 element in each step
  step_vector = 2.0 1

7.2.77.3 num_steps

  • Keywords Area
  • method
  • vector_parameter_study
  • num_steps

Number of sampling steps along the vector in a vector parameter study

Specification

Alias: none

Argument(s): INTEGER

Description

num_steps defines the number of steps that are taken in the direction of the vector. The magnitude of each step is determined in conjunction with the rest of the method specification.

Default Behavior

The user is required to specify num_steps for a vector parameter study. There is no default value.

This study performs function evaluations at both ends, making the total number of evaluations equal to num_steps+1.

Usage Tips

The study has stringent requirements on performing appropriate steps with any discrete range and discrete set variables. A num_steps specification must result in discrete range and set index steps that are integers: no remainder is currently permitted in the integer step calculation and no rounding to integer steps will occur.

Examples

method
  vector_parameter_study
  num_steps = 10
  final_point = 2.0 2.0

7.2.77.4 model_pointer

  • Keywords Area
  • method
  • vector_parameter_study
  • model_pointer

Identifier for model block to be used by a method
Keywords Area

Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
        0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
    surrogate global,
    dace_method_pointer = 'DACE'
    polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
    single
    interface_pointer = 'I1'
```
variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = ‘x1’ ‘x2’

interface
id_interface = ‘I1’
system async evaluation_concurrency = 5
analysis_driver = ‘text_book’

responses
response_functions = 3
no_gradients
no_hessians

7.2.78 list_parameter_study

  • Keywords Area
  • method
  • list_parameter_study

Samples variables as a specified values

Topics

This keyword is related to the topics:

  • parameter_studies

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Points Source (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>list_of_points</td>
<td>List of variable values to evaluate in a list parameter study</td>
</tr>
<tr>
<td></td>
<td></td>
<td>import_points_file</td>
<td>File containing list of variable values to evaluate in a list parameter study</td>
</tr>
</tbody>
</table>
Description

Dakota’s list parameter study allows for evaluations at user selected points of interest.

Default Behavior

By default, the list parameter study operates over all types of variables.

The number of real values in the list_of_points specification or file referenced by import_points_file must be a multiple of the total number of variables (including continuous and discrete types) contained in the variables specification.

Expected Outputs

A list parameter study produces a set of responses for each parameter set that is specified.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Parameter Sets

Usage Tips

- This parameter study simply performs simulations for the first parameter set (the first \( n \) entries in the list), followed by the next parameter set (the next \( n \) entries), and so on, until the list of points has been exhausted.

- Since the initial values from the variables specification will not be used, they need not be specified.

- When the points are specified in the Dakota input file using the list_of_points keyword, discrete set values must be referred to using 0-based indexes, and not the values themselves. However, when using import_points_file, refer to discrete set values directly, not by index.

- For both list_of_points and import_points_file, Dakota expects the values for each evaluation to be ordered by type. The type ordering matches that of the variables section of this Reference Manual and of the listing in the Parameters file format section of the Dakota User’s Manual[5]. When multiple variables are present for a single type, the ordering within that type must match the order specified by the user in the variables section of her input file.

Examples

This shows the method and variables block of a Dakota input file that runs a list_parameter_study.

```
method list_parameter_study
  list_of_points =
    3.1e6  0.0029  0.31
    3.2e6  0.0028  0.32
    3.3e6  0.0027  0.34
    3.3e6  0.0026  0.36

variables
  continuous_design = 3
  descriptors = ‘E’ ’MASS’ ’DENSITY’
```

Note that because of the way Dakota treats whitespace, the above example is equivalent to:
method
  list_parameter_study
  list_of_points =
  3.1e6  0.0029  0.31  3.2e6  0.0028
  0.32   3.3e6  0.0027
  0.34   3.3e6  0.0026  0.36

variables
  continuous_design = 3
  descriptors = 'E' 'MASS' 'DENSITY'

Although the first example is much more readable.

And here’s a full input file:

# tested on Dakota 6.0 on 140501
environment
  tabular_data
    tabular_data_file 'List_param_study.dat'
method
  list_parameter_study
  list_of_points =
    0.1  0.1
    0.2  0.1
    0.3  0.0
    0.3  1.0

model
  single

variables
  active_design
    continuous_design = 2
    descriptors = 'x1' 'x2'
  continuous_state = 1
    descriptors = 'constant1'
    initial_state = 100

interface
  analysis_drivers 'text_book'
  fork
    asynchronous
      evaluation_concurrency 2

responses
  response_functions = 1
  no_gradients
  no_hessians

This example illustrates the list_parameter_study.

- The function evaluations are independent, so any level of evaluation_concurrency can be used
- Default behavior for parameter studies is to iterate on all variables. However, because active design is specified, this study will only iterate on the continuous_design variables.

See Also

These keywords may also be of interest:

- centered_parameter_study
- multidim_parameter_study
- vector_parameter_study
7.2.78.1 list_of_points

- Keywords Area

- method

- list_parameter_study

- list_of_points

List of variable values to evaluate in a list parameter study

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Description**

The `list_of_points` keyword allows the user to specify, in a freeform format, a list of variable values at which to compute a model response.

**Default Behavior**

The user is required to provide a list of points for a list parameter study either by specifying it with `list_of_points` or by providing a file from which such a list can be read via `import_points_file`. There is no default list of points.

**Usage Tips**

The number of values in the list must be an integer multiple of the number of variables. Dakota will verify that this condition is met.

**Examples**

```plaintext
method
list_parameter_study
list_of_points =
  3.1e6  0.0029  0.31
  3.2e6  0.0028  0.32
  3.3e6  0.0027  0.34
  3.3e6  0.0026  0.36
```

7.2.78.2 import_points_file

- Keywords Area

- method

- list_parameter_study

- import_points_file

File containing list of variable values to evaluate in a list parameter study
7.2 method

Specification

Alias: none

Argument(s): STRING

Default: no point import from a file

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td></td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

Description

The import_points_file specifies a file containing a list of variable values at which to compute a model response.

Default Behavior

By default, methods do not import points from a file.

Usage Tips

Dakota parses input files without regard to whitespace, but the import_points_file must be in one of three formats:

- annotated (default)
- custom_annotated
- freeform

Examples

```dakota
method
  list_parameter_study
    import_points_file = 'dakota_pstudy.3.dat'
```

7.2.78.2.1 custom_annotated

- Keywords Area
- method
- list_parameter_study
- import_points_file
- custom_annotated

Selects custom-annotated tabular file format
Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

**Examples**

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:
7.2 method

environment
taxable_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

7.2.78.2.1.1 header

- Keywords Area
- method
- list_parameter_study
- import_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated

7.2.78.2.1.2 eval_id

- Keywords Area
- method
- list_parameter_study
- import_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Description

See description of parent `custom_annotated`

7.2.78.2.1.3  `interface_id`

- Keywords Area
- `method`
- `list_parameter_study`
- `import_points_file`
- `custom_annotated`
- `interface_id`

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent `custom_annotated`

7.2.78.2.2  `annotated`

- Keywords Area
- `method`
- `list_parameter_study`
- `import_points_file`
- `annotated`

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- `file_formats`

Specification

Alias: none
Argument(s): none
Default: annotated
Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated

Resulting tabular file:
```

```
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.2method freeform

- Keywords Area

- method

- list_parameter_study

- import_points_file

- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Specification

**Alias:** none  
**Argument(s):** none

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The *annotated* format is the default for tabular export/import. To change this behavior, specify *freeform* in the relevant export/import context.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to *annotated* format, though *freeform* remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In *freeform*, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

**Examples**

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform

Resulting tabular file:

```
0.9  1.1  0.0002  0.26  0.76  
0.90009  1.1  0.0001996404857  0.2601620081  0.759955  
0.89991  1.1  0.0002003604863  0.2598380081  0.760045  
...
```

7.2.78.2.4  active_only

- **Keywords Area**
- **method**
- **list_parameter_study**
- **import_points_file**
- **active_only**

Import only active variables from tabular data file
7.2 method

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.2.78.3 model_pointer

- Keywords Area
- method
- list_parameter_study
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.
Usage Tips
When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = ‘UQ’

method
  id_method = ‘UQ’
  model_pointer = ‘SURR’
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
                    0.1 0.2 0.6
                    0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = ‘SURR’
  surrogate global,
  dace_method_pointer = ‘DACE’
  polynomial quadratic

method
  id_method = ‘DACE’
  model_pointer = ‘DACE_M’
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = ‘DACE_M’
  single
  interface_pointer = ‘I1’

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = ‘x1’ ‘x2’

interface
  id_interface = ‘I1’
  system asynch evaluation_concurrency = 5
  analysis_driver = ‘text_book’

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.79 centered_parameter_study

- Keywords Area
- method
- centered_parameter_study

Samples variables along points moving out from a center point
Topics

This keyword is related to the topics:

- parameter_studies

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td></td>
<td>step_vector</td>
<td>Size of steps to be taken in each dimension of a centered parameter study</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>steps_per_variable</td>
<td>Number of steps to take in each dimension of a centered parameter study</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>model_pointer</td>
<td>Identifier for model block to be used by a method</td>
</tr>
</tbody>
</table>

Description

Dakota’s centered parameter study computes response data sets along multiple coordinate-based vectors, one per parameter, centered about the initial values from the variables specification. This is useful for investigation of function contours with respect to each parameter individually in the vicinity of a specific point (e.g., post-optimality analysis for verification of a minimum), thereby avoiding the cost associated with a multidimensional grid.

**Default Behavior**

By default, the centered parameter study operates over all types of variables.

The centered_parameter_study takes steps along each orthogonal dimension. Each dimension is treated independently. The number of steps are taken in each direction, so that the total number of points in the parameter study is $1 + 2 \sum n$.

**Expected Outputs**

A centered parameter study produces a set of responses for each parameter set that is generated.

**Expected HDF5 Output**

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Parameter Sets
- Variable Slices
Examples

The following example is a good comparison to the examples on `multidim_parameter_study` and `vector_parameter_study`.

```bash
# tested on Dakota 6.0 on 140501
environment
tabular_data
tabular_data_file = 'rosen_centered.dat'

method
centered_parameter_study
  steps_per_variable = 5 4
  step_vector = 0.4 0.5

model
  single

variables
  continuous_design = 2
  initial_point = 0 0
  descriptors = 'x1' "x2"

interface
  analysis_driver = 'rosenbrock'
  fork

responses
  response_functions = 1
  no_gradients
  no_hessians
```

See Also

These keywords may also be of interest:

- `multidim_parameter_study`
- `list_parameter_study`
- `vector_parameter_study`

7.2.79.1 step_vector

- Keywords Area
- method
- centered_parameter_study
- step_vector

Size of steps to be taken in each dimension of a centered parameter study

Specification

**Alias:** none

**Argument(s):** REALLIST
Description

The `step_vector` keyword defines the individual step size in each dimension, treated separately.

Default Behavior

The user is required to define the number of step sizes for a centered parameter study. There are no default values.

Steps are taken in the plus and minus directions, and are defined in either actual values (continuous and discrete range) or index offsets (discrete set).

Examples

```plaintext
method centered_parameter_study
   steps_per_variable = 5 4
   step_vector = 0.4 0.5
```

7.2.79.2 steps_per_variable

- Keywords Area
- method
- centered_parameter_study
- steps_per_variable

Number of steps to take in each dimension of a centered parameter study

Specification

Alias: `deltas_per_variable`

Argument(s): INTEGERLIST

Description

The `steps_per_variable` keyword allows the user to define the number of steps in each dimension of a centered parameter study. Because they are taken independently, the number of steps can be specified for each.

Default Behavior

The user is required to define the number of steps per variable for a centered parameter study. There are no default values.

Steps are taken in the plus and minus directions, and are defined in either actual values (continuous and discrete range) or index offsets (discrete set).

Examples

```plaintext
method centered_parameter_study
   steps_per_variable = 5 4
   step_vector = 0.4 0.5
```
7.2.79.3 model_pointer

- Keywords Area
- method
- centered_parameter_study
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
      samples = 10
      seed = 98765 rng rnum2
      response_levels = 0.1 0.2 0.6
      0.1 0.2 0.6
      0.1 0.2 0.6
  sample_type lhs
  distribution cumulative

model
  id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
  samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
  lower_bounds = 0. 0.
  upper_bounds = 1. 1.
  descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
  analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.80 multidim_parameter_study

  • Keywords Area
  • method
  • multidim_parameter_study

Samples variables on full factorial grid of study points

Topics

This keyword is related to the topics:

  • parameter_studies

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>partitions</td>
<td></td>
<td>Samples variables on full factorial grid of study points</td>
</tr>
</tbody>
</table>
Description

Dakota’s multidimensional parameter study computes response data sets for an n-dimensional grid of points. Each continuous and discrete range variable is partitioned into equally spaced intervals between its upper and lower bounds, each discrete set variable is partitioned into equally spaced index intervals. The partition boundaries in n-dimensional space define a grid of points, and every point is evaluated.

Default Behavior

By default, the multidimensional parameter study operates over all types of variables.

Expected Outputs

A multidimensional parameter study produces a set of responses for each parameter set that is generated.

Expected HDF5 Output

If Dakota was built with HDF5 support and run with the hdf5 keyword, this method writes the following results to HDF5:

- Parameter Sets
- Correlations

Usage Tips

Since the initial values from the variables specification will not be used, they need not be specified.

Examples

This example is taken from the Users Manual and is a good comparison to the examples on centered_parameter_study and vector_parameter_study.

```bash
# tested on Dakota 6.0 on 140501
environment
  tabular_data
    tabular_data_file = 'rosen_multidim.dat'

method
  multidim_parameter_study
  partitions = 10 8

model
  single

variables
  continuous_design = 2
    lower_bounds -2.0 -2.0
    upper_bounds 2.0 2.0
  descriptors 'x1' "x2"

interface
  analysis_driver = 'rosenbrock'
  fork

responses
  response_functions = 1
  no_gradients
  no_hessians
```

This example illustrates the full factorial combinations of parameter values created by the multidim_parameter_study. With 10 and 8 partitions, there are actually 11 and 9 values for each variable. This means that $11 \times 9 = 99$
function evaluations will be required.

See Also

These keywords may also be of interest:

- centered_parameter_study
- list_parameter_study
- vector_parameter_study

7.2.80.1 partitions

- Keywords Area
- method
- multidim_parameter_study
- partitions

Samples variables on full factorial grid of study points

Topics

This keyword is related to the topics:

- parameter_studies

Specification

Alias: none  
Argument(s): INTEGERLIST

Description

Dakota's multidimensional parameter study computes response data sets for an n-dimensional grid of points. Each continuous and discrete range variable is partitioned into equally spaced intervals between its upper and lower bounds, each discrete set variable is partitioned into equally spaced index intervals. The partition boundaries in n-dimensional space define a grid of points, and every point is evaluated.

Default Behavior

By default, the multidimensional parameter study operates over all types of variables.

Expected Outputs

A multidimensional parameter study produces a set of responses for each parameter set that is generated.

Usage Tips

Since the initial values from the variables specification will not be used, they need not be specified.
Examples

This example is taken from the Users Manual and is a good comparison to the examples on centered_parameter_study and vector_parameter_study.

```plaintext
# tested on Dakota 6.0 on 140501
environment
tabular_data
tabular_data_file = 'rosen_multidim.dat'

method
multidim_parameter_study
partitions = 10 8

model
single

variables
  continuous_design = 2
    lower_bounds -2.0 -2.0
    upper_bounds 2.0 2.0
    descriptors 'x1' "x2"

interface
  analysis_driver = 'rosenbrock'
  fork

responses
  response_functions = 1
  no_gradients
  no_hessians
```

This example illustrates the full factorial combinations of parameter values created by the multidim_parameter_study. With 10 and 8 partitions, there are actually 11 and 9 values for each variable. This means that $11 \times 9 = 99$ function evaluations will be required.

See Also

These keywords may also be of interest:

- centered_parameter_study
- list_parameter_study
- vector_parameter_study

7.2.80.2 model_pointer

- Keywords Area
- method
- multidim_parameter_study
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer
Specification

**Alias:** none

**Argument(s):** STRING

**Default:** method use of last model parsed (or use of default model if none parsed)

Description

The `model_pointer` is used to specify which `model` block will be used to perform the function evaluations needed by the Dakota method.

**Default Behavior**

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a `model` block in the Dakota input file that has a corresponding `id_model` with the same name.

**Usage Tips**

When doing advanced analyses that involve using multiple methods and multiple models, defining a `model_pointer` for each method is imperative.

See `block_pointer` for details about pointers.

Examples

```
environment
tabular_data
method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
    samples = 10
    seed = 98765 rng rnum2
    response_levels = 0.1 0.2 0.6
        0.1 0.2 0.6
        0.1 0.2 0.6
    sample_type lhs
    distribution cumulative

model
  id_model = 'SURR'
  surrogate global,
  dace_method_pointer = 'DACE'
  polynomial quadratic

method
  id_method = 'DACE'
  model_pointer = 'DACE_M'
  sampling sample_type lhs
    samples = 121 seed = 5034 rng rnum2

model
  id_model = 'DACE_M'
  single
  interface_pointer = 'I1'

variables
  uniform_uncertain = 2
    lower_bounds = 0. 0.
    upper_bounds = 1. 1.
    descriptors = 'x1' 'x2'

interface
  id_interface = 'I1'
  system asynch evaluation_concurrency = 5
```
analysis_driver = 'text_book'

responses
  response_functions = 3
  no_gradients
  no_hessians

7.2.81 richardson_extrap

- Keywords Area
- method
- richardson_extrap

Estimate order of convergence of a response as model fidelity increases

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Verification Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>estimate_order</td>
<td>Compute the best estimate of the convergence order from three points</td>
</tr>
<tr>
<td></td>
<td></td>
<td>converge_order</td>
<td>Refine until the estimated convergence order converges</td>
</tr>
<tr>
<td></td>
<td></td>
<td>converge_qoi</td>
<td>Refine until the response converges</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>refinement_rate</td>
<td>Rate at which the state variables are refined</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>convergence_-tolerance</td>
<td>Stopping criterion based on objective function or statistics convergence</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_iterations</td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
</tbody>
</table>
Solution verification procedures estimate the order of convergence of the simulation response data during the course of a refinement study. This branch of methods is new and currently only contains one algorithm: Richardson extrapolation.

Refinement of the model

The model fidelity must be parameterized by one or more continuous state variable(s).

The refinement path is determined from the initial state of the continuous state variables specification in combination with the refinement rate, where each of the state variables is treated as an independent refinement factor and each of the initial state values is repeatedly divided by the refinement rate value to define new discretization states.

Results

Three algorithm options are currently provided:

1. estimate_order
2. converge_order
3. converge_qoi

Stopping Criteria

The method employs the max_iterations and convergence_tolerance method independent controls as stopping criteria.

Theory

In each of these cases, convergence order for a response quantity of interest (QoI) is estimated from

\[ p = \ln \left( \frac{QoI_3 - QoI_2}{QoI_2 - QoI_1} \right) / \ln(r) \]

where \( r \) is the uniform refinement rate specified by refinement_rate.

7.2.81.1 estimate_order

- Keywords Area
- method
- richardson_extrap
- estimate_order

Compute the best estimate of the convergence order from three points

Specification

Alias: none

Argument(s): none
The **estimate_order** option is the simplest option. For each of the refinement factors, it evaluates three points along the refinement path and uses these results to perform an estimate of the convergence order for each response function.

### 7.2.81.2 **converge_order**

- **Keywords Area**
- **method**
- **richardson_extrap**
- **converge_order**

Refine until the estimated convergence order converges

### Specification

**Alias:** none  
**Argument(s):** none

### Description

The **converge_order** option is initialized using the **estimate_order** approach, and additional refinements are performed along the refinement path until the convergence order estimates converge (two-norm of the change in response orders is less than the convergence tolerance).

### 7.2.81.3 **converge_qoi**

- **Keywords Area**
- **method**
- **richardson_extrap**
- **converge_qoi**

Refine until the response converges

### Specification

**Alias:** none  
**Argument(s):** none

### Description

The **converge_qoi** option is similar to the **converge_order** option, except that the convergence criterion is that the two-norm of the response discretization errors (computed from extrapolation) must be less than the convergence tolerance.
7.2.81.4 refinement_rate

Keyword Area

- method
- richardson_extrap
- refinement_rate

Rate at which the state variables are refined

Specification

Alias: none
Argument(s): REAL
Default: 2.

Description

Described on parent page

7.2.81.5 convergence_tolerance

Keyword Area

- method
- richardson_extrap
- convergence_tolerance

Stopping criterion based on objective function or statistics convergence

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

Alias: none
Argument(s): REAL
Default: 1.e-4

Description

The convergence_tolerance specification provides a real value for controlling the termination of iteration. For optimization, it is most commonly a relative convergence tolerance for the objective function; i.e., if the change in the objective function between successive iterations divided by the previous objective function is less than the amount specified by convergence_tolerance, then this convergence criterion is satisfied on the current iteration.
Therefore, permissible values are between 0 and 1, non-inclusive.

**Behavior Varies by Package/Library**

This control is used with most optimization and least squares iterators (DOT, CONMIN, NLPQLP, NPSOL, NLSSOL, OPT++, and SCOLIB). Most other Dakota methods (such as DACE or parameter studies) do not use this control, but some adaptive methods, such as adaptive UQ, do.

Since no progress may be made on one iteration followed by significant progress on a subsequent iteration, some libraries require that the convergence tolerance be satisfied on two or more consecutive iterations prior to termination of iteration.

Notes on each library:

- DOT: relative tolerance that must be satisfied for two consecutive iterations
- NL2SOL: See nl2sol
- NLPQLP: used as Lagrangian gradient norm tolerance (ACC), not as a relative convergence tolerance
- NPSOL: used as a line search tolerance, not as a relative convergence tolerance

7.2.81.6 max_iterations

- **Keywords Area**
- **method**
- **richardson_extrap**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

**Topics**

This keyword is related to the topics:

- method_independent_controls

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100 (exceptions: fsu_cvt, local_reliability: 25; global_(reliability, interval_est, evidence) / efficient_global: 25*n)

**Description**

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed \texttt{max_iterations} iterations. In cases where there are multiple nested iterations to be controlled, \texttt{max_iterations} generally controls the outer-most context.

**Default Behavior**

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.2.81.7 model_pointer

- Keywords Area
- method
- richardson_extrap
- model_pointer

Identifier for model block to be used by a method

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed (or use of default model if none parsed)

Description

The model_pointer is used to specify which model block will be used to perform the function evaluations needed by the Dakota method.

Default Behavior

If not specified, a Dakota method will use the last model block parsed. If specified, there must be a model block in the Dakota input file that has a corresponding id_model with the same name.

Usage Tips

When doing advanced analyses that involve using multiple methods and multiple models, defining a model_pointer for each method is imperative.

See block_pointer for details about pointers.

Examples

```plaintext
environment
  tabular_data
  method_pointer = 'UQ'

method
  id_method = 'UQ'
  model_pointer = 'SURR'
  sampling,
  samples = 10
```
seed = 98765 rng rnum2
response_levels = 0.1 0.2 0.6
                  0.1 0.2 0.6
                  0.1 0.2 0.6
sample_type lhs
distribution cumulative

model
id_model = 'SURR'
surrogate global,
dace_method_pointer = 'DACE'
polynomial quadratic

method
id_method = 'DACE'
model_pointer = 'DACE_M'
sampling sample_type lhs
samples = 121 seed = 5034 rng rnum2

model
id_model = 'DACE_M'
single
interface_pointer = 'I1'

variables
uniform_uncertain = 2
lower_bounds = 0. 0.
upper_bounds = 1. 1.
descriptors = 'x1' 'x2'

interface
id_interface = 'I1'
system asynch evaluation_concurrency = 5
analysis_driver = 'text_book'

responses
response_functions = 3
no_gradients
no_hessians

7.3 model

• Keywords Area

• model

Specifies how variables are mapped into a set of responses

Topics

This keyword is related to the topics:

• block

Specification

Alias: none

Argument(s): none

Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>id_model</td>
<td>Give the model block an identifying name, in case of multiple model blocks</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Model Type (Group 1)</td>
<td>single</td>
<td>A model with one of each block: variable, interface, and response</td>
</tr>
<tr>
<td></td>
<td></td>
<td>surrogate</td>
<td>An empirical model that is created from data or the results of a submodel</td>
</tr>
<tr>
<td></td>
<td></td>
<td>nested</td>
<td>A model whose responses are computed through the use of a sub-iterator</td>
</tr>
<tr>
<td></td>
<td></td>
<td>active_subspace</td>
<td>Active (variable) subspace model</td>
</tr>
<tr>
<td></td>
<td></td>
<td>adapted_basis</td>
<td>Basis adaptation model</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random_field</td>
<td>Experimental capability to generate a random field representation. From data, from simulation runs, or from a covariance matrix. The representation may then be sampled for use as a random field input to another simulation. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
</tbody>
</table>
Description

A model is comprised of a mapping from variables, through an interface, to responses.

Model Group 1 The type of model can be:

1. single (default)
2. surrogate
3. nested
4. subspace
5. random_field

The input file must specify one of these types. If the type is not specified, Dakota will assume a single model.

Block Pointers and ID

Each of these model types supports `variables_pointer` and `responses_pointer` strings for identifying the variables and responses specifications used in constructing the model by cross-referencing with `id_variables` and `id_responses` strings from particular variables and responses keyword specifications.

These pointers are valid for each model type since each model contains a set of variables that is mapped into a set of responses – only the specifics of the mapping differ.

Additional pointers are used for each model type for constructing the components of the variable to response mapping. As an environment specification identifies a top-level method and a method specification identifies a model, a model specification identifies variables, responses, and (for some types) interface specifications. This top-down flow specifies all of the object interrelationships.

Examples

The first example shows a minimal specification for a single model, which is the default model when no models are explicitly specified by the user.

```dakota
model
  single
```

The next example displays a surrogate model specification which selects a quadratic polynomial from among the global approximation methods. It uses a pointer to a design of experiments method for generating the data needed for building the global approximation, reuses any old data available for the current approximation region, and employs the first-order multiplicative approach to correcting the approximation each time correction is requested.

```dakota
model,
  id_model = 'M1'
  variables_pointer = 'V1'
  responses_pointer = 'R1'
```
surrogate
global
    polynomial quadratic
dace_method_pointer = 'DACE'
reuse_samples region
correction multiplicative first_order

This example demonstrates the use of identifiers and pointers. It provides the optional model independent specifications for model identifier, variables pointer, and responses pointer as well as model dependent specifications for global surrogates (see global).

Finally, an advanced nested model example would be

model
    id_model = 'M1'
    variables_pointer = 'V1'
    responses_pointer = 'R1'
nested
    optional_interface_pointer = 'OI1'
    optional_interface_responses_pointer = 'OIR1'
    sub_method_pointer = 'SM1'
    primary_variable_mapping = '' 'X' 'Y'
    secondary_variable_mapping = '' 'mean' 'mean'
    primary_response_mapping = 1. 0. 0. 0. 0. 0. 0. 0. 0.
    secondary_response_mapping = 0. 0. 0. 1. 3. 0. 0. 0. 0.
    0. 0. 0. 0. 0. 1. 3. 0.

This example illustrates controls for model identifier, variables pointer, and responses pointer and for specifying details of the nested mapping.

7.3.1 id_model

- Keywords Area
- model
- id_model

Give the model block an identifying name, in case of multiple model blocks

Topics

This keyword is related to the topics:

- block_identifier

Specification

Alias: none

Argument(s): STRING

Default: method use of last model parsed

Description

The model identifier string is supplied with id_model and is used to provide a unique identifier string for use within method specifications (refer to the keyword model_pointer in any of the methods in the method block, for example: model_pointer)

This is used to determine which model the method will run.
See Also

These keywords may also be of interest:

- model_pointer

### 7.3.2 single

- Keywords Area
- model
- single

A model with one of each block: variable, interface, and response

**Specification**

**Alias:** simulation

**Argument(s):** none

**Default:** N/A ( single if no model specification)

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>interface_pointer</td>
<td>Interface block pointer for the single model type</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>solution_level_cost</td>
<td>Cost estimates associated with a set of solution control values.</td>
</tr>
</tbody>
</table>

**Description**

The single model is the simplest model type. It uses a single interface instance to map variables into responses. There is no recursion in this case.

The optional interface_pointer specification identifies the interface block by cross-referencing with the id_interface string input from a particular interface keyword specification. This is only necessary when the input file has multiple interface blocks, and you wish to explicitly point to the desired block. The same logic follows for responses and variables blocks and pointers.

**Examples**

The example shows a minimal specification for a single model, which is the default model when no models are specified by the user.

```plaintext
model
single
```

This example does not provide any pointer strings and therefore relies on the default behavior of constructing the model with the last variables, interface, and responses specifications parsed.
7.3 model

See Also

These keywords may also be of interest:

- surrogate
- nested

7.3.2.1 interface_pointer

- Keywords Area
- model
- single
- interface_pointer

Interface block pointer for the single model type

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none
Argument(s): STRING
Default: model use of last interface parsed

Description

In the single model case, a single interface is used to map the variables into responses. The optional interface_pointer specification identifies this interface by cross-referencing with the id_interface string input from a particular interface keyword specification.

See block_pointer for details about pointers.

When an interface pointer is not specified, the model will use the last interface block parsed from the input file.

7.3.2.2 solution_level_cost

- Keywords Area
- model
- single
- solution_level_cost

Cost estimates associated with a set of solution control values.
Specification

**Alias:** none

**Argument(s):** REALLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>solution_level_control</td>
<td>Cost estimates associated with a set of solution control values.</td>
</tr>
</tbody>
</table>

Description

Simulation-based models may have an associated `solution_level_control`, which identifies a hierarchy of solution states, such as a set of mesh discretizations from coarse to fine, a set of solver tolerances from loose to tight, etc. In algorithms that manage such a hierarchy and perform optimal resource allocation among the solution states (e.g., multilevel Monte Carlo), it is important to estimate a set of costs associated with each state. These cost estimates can be relative, such as in the example below (lowest cost normalized to 1.)

**Note:** a scalar solution cost can be specified without an associated solution level control. This is useful when employing a hierarchy of model forms (each model has a scalar solution cost and no solution level control) instead of a hierarchy of discretization levels (one model has a vector-valued solution cost associated with multiple solution levels).

Examples

```plaintext
model,
  simulation
    solution_level_control = 'mesh_size'
    solution_level_cost = 1. 8. 64. 512. 4096.
variables,
  uniform_uncertain = 9
  lower_bounds = 9*-1.
  upper_bounds = 9* 1.
  discrete_state_set
    integer = 1
      set_values = 4 8 16 32 64
      descriptors = 'mesh_size'
```

7.3.2.2.1 `solution_level_control`

- **Keywords Area**
- **model**
- **single**
- **solution_level_cost**
- **solution_level_control**

Cost estimates associated with a set of solution control values.

Specification

**Alias:** none
Argument(s): STRING

Default: use of single default solution level

Description

Simulation-based models may have an associated solution_level_control, which identifies a hierarchy of solution states, such as a set of mesh discretizations from coarse to fine, a set of solver tolerances from loose to tight, etc. The string specified as the solution_level_control identifies a discrete variable label that parameterizes the hierarchy of solution states.

Note: If the discrete variable identified is a discrete set variable, then it is important to note that the variable’s set values will be ordered (lexicographically in the case of string variables), and the ordering of values provided in solution_level_cost should correspond to this set ordering. A common error is to provide a listing of set values that is out of order and then providing a set of costs corresponding to this misordered list – in this case, the solution level costs will be associated with the re-ordered set values.

Examples

In this example, integer solution control values and solution costs are naturally well ordered.

```plaintext
model,
  simulation
      solution_level_control = 'mesh_size'
      solution_level_cost = 1. 8. 64. 512. 4096.
variables,
  uniform_uncertain = 9
  lower_bounds     = 9*-1.
  upper_bounds     = 9* 1.
  discrete_state_set
    integer = 1
    set_values = 4 8 16 32 64
    descriptors = 'mesh_size'
```

In this example, string solution control values are lexicographically ordered, and care must be taken to align the solution cost estimates.

```plaintext
model,
  simulation
      solution_level_control = 'mesh_size'
      solution_level_cost = 1. 64. 8. # match set ordering
variables,
  uniform_uncertain = 9
  lower_bounds     = 9*-1.
  upper_bounds     = 9* 1.
  discrete_state_set
    string = 1
    set_values = 'COARSE' 'FINE' 'MEDIUM' # lexicographical ordering
    descriptors = 'mesh_size'
```

7.3.3 surrogate

- Keywords Area
- model
- surrogate

An empirical model that is created from data or the results of a submodel
### Specification

**Alias:** none  
**Argument(s):** none  

### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>id_surrogates</td>
<td>Identifies the subset of the response functions by number that are to be approximated (the default is all functions). Select a surrogate model with global support</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Surrogate Category (Group 1)</td>
<td>global</td>
<td></td>
</tr>
<tr>
<td>multipoint</td>
<td>Build a locally accurate surrogate from data at a single point</td>
<td></td>
<td></td>
</tr>
<tr>
<td>local</td>
<td>Construct a surrogate from multiple existing training points</td>
<td></td>
<td></td>
</tr>
<tr>
<td>hierarchical</td>
<td>Build a locally accurate surrogate from data at a single point</td>
<td></td>
<td></td>
</tr>
<tr>
<td>non_hierarchical</td>
<td>Hierarchical surrogates use an ordered hierarchy of model fidelities and/or resolutions to trade accuracy versus cost.</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Description

Surrogate models are inexpensive approximate models that are intended to capture the salient features of an expensive high-fidelity model. They can be used to explore the variations in response quantities over regions of the parameter space, or they can serve as inexpensive stand-ins for optimization or uncertainty quantification studies (see, for example, the surrogate-based optimization methods, surrogate_based_global and surrogate_based_local).

Surrogate models supported in Dakota are categorized as Data Fitting or Hierarchical, as shown below. Each of these surrogate types provides an approximate representation of a “truth” model which is used to perform the parameter to response mappings. This approximation is built and updated using results from the truth model, called the “training data”.

- Data fits:
Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

Data fitting methods involve construction of an approximation or surrogate model using data (response values, gradients, and Hessians) generated from the original truth model. Data fit methods can be further categorized as local, multipoint, and global approximation techniques, based on the number of points used in generating the data fit.

1. Local: built from response data from a single point in parameter space
   - Taylor series expansion: taylor_series
     Training data consists of a single point, plus gradient and Hessian information.

2. Multipoint: built from two or more points in parameter space, often involving the current and previous iterates of a minimization algorithm.
   - TANA-3: tana
     Training Data comes from a few previously evaluated points

3. Global full space response surface methods:
   - Polynomial regression: polynomial
   - Gaussian process (Kriging): gaussian_process
   - Artificial neural network: neural_network
   - MARS: mars
   - Radial Basis Functions: radial_basis
   - Orthogonal polynomials (only supported in PCE/SC for now): polynomial_chaos and stoch_collocation
     Training data is generated using either a design of experiments method applied to the truth model (specified by dace_method_pointer), or from saved data (specified by reuse_points) in a restart database, or an import file.

   • Multifidelity/hierarchical:
     Multifidelity modeling involves the use of a low-fidelity physics-based model as a surrogate for the original high-fidelity model. The low-fidelity model typically involves a coarser mesh, looser convergence tolerances, reduced element order, or omitted physics.
     See hierarchical.

The global and hierarchal surrogates have a correction feature in order to improve the local accuracy of the surrogate models. The correction factors force the surrogate models to match the true function values and possibly true function derivatives at the center point of each trust region. Details can be found on global correction or hierarchical correction.

Theory

Surrogate models are used extensively in the surrogate-based optimization and least squares methods, in which the goals are to reduce expense by minimizing the number of truth function evaluations and to smooth out noisy data with a global data fit. However, the use of surrogate models is not restricted to optimization techniques; uncertainty quantification and optimization under uncertainty methods are other primary users.

Data Fit Surrogate Models

A surrogate of the [data fit] type is a non-physics-based approximation typically involving interpolation or regression of a set of data generated from the original model. Data fit surrogates can be further characterized by the number of data points used in the fit, where a local approximation (e.g., first or second-order Taylor series) uses data from a single point, a multipoint approximation (e.g., two-point exponential approximations (TPEA) or two-point adaptive nonlinearity approximations (TANA)) uses a small number of data points often drawn from the previous iterates of a particular algorithm, and a global approximation (e.g., polynomial response surfaces, kriging/gaussian_process,
neural networks, radial basis functions, splines) uses a set of data points distributed over the domain of interest, often generated using a design of computer experiments.

Dakota contains several types of surface fitting methods that can be used with optimization and uncertainty quantification methods and strategies such as surrogate-based optimization and optimization under uncertainty. These are: polynomial models (linear, quadratic, and cubic), first-order Taylor series expansion, kriging spatial interpolation, artificial neural networks, multivariate adaptive regression splines, radial basis functions, and moving least squares. With the exception of Taylor series methods, all of the above methods listed in the previous sentence are accessed in Dakota through the Surfpack library. All of these surface fitting methods can be applied to problems having an arbitrary number of design parameters. However, surface fitting methods usually are practical only for problems where there are a small number of parameters (e.g., a maximum of somewhere in the range of 30-50 design parameters). The mathematical models created by surface fitting methods have a variety of names in the engineering community. These include surrogate models, meta-models, approximation models, and response surfaces. For this manual, the terms surface fit model and surrogate model are used.

The data fitting methods in Dakota include software developed by Sandia researchers and by various researchers in the academic community.

**Multifidelity Surrogate Models**

A second type of surrogate is the \{model hierarchy\} type (also called multifidelity, variable fidelity, variable complexity, etc.). In this case, a model that is still physics-based but is of lower fidelity (e.g., coarser discretization, reduced element order, looser convergence tolerances, omitted physics) is used as the surrogate in place of the high-fidelity model. For example, an inviscid, incompressible Euler CFD model on a coarse discretization could be used as a low-fidelity surrogate for a high-fidelity Navier-Stokes model on a fine discretization.

**Surrogate Model Selection**

This section offers some guidance on choosing from among the available surrogate model types.

- For Surrogate Based Local Optimization, using the surrogate\_based\_local method with a trust region:
  - using the keywords:
    1. surrogate local taylor\_series or
    2. surrogate multipoint tana
  - will probably work best.
  If for some reason you wish or need to use a global surrogate (not recommended) then the best of these options is likely to be either:
    1. surrogate global gaussian\_process surfpack or
    2. surrogate global moving\_least\_squares.

- For Efficient Global Optimization (EGO), the efficient\_global method:
  - the default surrogate is: gaussian\_process surfpack which is likely to find a more optimal value and/or require fewer true function evaluations than the alternative, gaussian\_process dakota. However, the surfpack will likely take more time to build than the dakota version. Note that currently the use\_derivatives keyword is not recommended for use with EGO based methods.

- For EGO based global interval estimation, the global\_interval\_est ego method:
  - the default gaussian\_process surfpack will likely work better than the alternative gaussian\_process dakota.

- For Efficient Global Reliability Analysis (EGRA), the global\_reliability method:
  - the surfpack and dakota versions of the gaussian process tend to give similar answers with the dakota version tending to use fewer true function evaluations. Since this is based on EGO, it is likely that the default surfpack is more accurate, although this has not been rigorously demonstrated.

- For EGO based Dempster-Shafer Theory of Evidence, i.e. the global\_evidence ego method, the default gaussian\_process surfpack often use significantly fewer true function evaluations than the alternative gaussian\_process dakota.

- When using a global surrogate to extrapolate, any of the surrogates:
are recommended.

• When there is over roughly two or three thousand data points and you wish to interpolate (or approximately inter-polate) then a Taylor series, Radial Basis Function Network, or Moving Least Squares fit is recommended. The only reason that the \texttt{gaussian\_process surfpack} is not recommended is that it can take a considerable amount of time to construct when the number of data points is very large. Use of the third party MARS package included in Dakota is generally discouraged.

• In other situations that call for a global surrogate, the \texttt{gaussian\_process surfpack} is generally recommended. The \texttt{use\_derivatives} keyword will only be useful if accurate and inexpensive derivatives are available. Finite difference derivatives are disqualified on both counts. However, derivatives generated by analytical, automatic differentiation, or continuous adjoint techniques can be appropriate. Currently, first order derivatives, i.e. gradients, are the highest order derivatives that can be used to construct the \texttt{gaussian\_process surfpack} model; Hessians will not be used even if they are available.

See Also

These keywords may also be of interest:

• single

• nested

7.3.3.1 \texttt{id\_surrogates}

• Keywords Area

• model

• surrogate

• id\_surrogates

Identifies the subset of the response functions by number that are to be approximated (the default is all functions).

Specification

\textbf{Alias:} none

\textbf{Argument(s):} INTEGERLIST

\textbf{Default:} All response functions are approximated

Description

In the \texttt{surrogate} model case, the specification first allows a mixture of surrogate and actual response mappings through the use of the optional \texttt{id\_surrogates} specification. This identifies the subset of the response functions by number that are to be approximated (the default is all functions). The valid response function identifiers range from 1 through the total number of response functions (see \texttt{response\_functions}).
Select a surrogate model with global support

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Global Surrogate Type (Group 1)</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>Global Surrogate Type (Group 1)</td>
<td>Use the Gaussian process regression surrogate from the surrogates module</td>
</tr>
<tr>
<td><code>gaussian_process</code></td>
<td><code>gaussian_process</code></td>
<td>Gaussian Process surrogate model</td>
</tr>
<tr>
<td><code>mars</code></td>
<td><code>mars</code></td>
<td>Multivariate Adaptive Regression Spline (MARS)</td>
</tr>
<tr>
<td><code>moving_least_squares</code></td>
<td><code>moving_least_squares</code></td>
<td>Moving Least Squares surrogate models</td>
</tr>
<tr>
<td><code>function_train</code></td>
<td><code>function_train</code></td>
<td>Global surrogate model based on functional tensor train decomposition</td>
</tr>
<tr>
<td><code>neural_network</code></td>
<td><code>neural_network</code></td>
<td>Artificial neural network model</td>
</tr>
<tr>
<td><code>radial_basis</code></td>
<td><code>radial_basis</code></td>
<td>Radial basis function (RBF) model</td>
</tr>
<tr>
<td>Optional</td>
<td>Number of Build Points (Group 2)</td>
<td>Polynomial surrogate model</td>
</tr>
<tr>
<td>----------</td>
<td>---------------------------------</td>
<td>-----------------------------</td>
</tr>
<tr>
<td></td>
<td>total_points</td>
<td>Use a deterministic polynomial surrogate</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>domain_decomposition</td>
<td>Piecewise Domain Decomposition for Global Surrogate Models</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>total_points</td>
<td>Specified number of training points</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>minimum_points</td>
<td>Construct surrogate with minimum number of points</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>recommended_points</td>
<td>Construct surrogate with recommended number of points</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Build Data Source (Group 3)</td>
<td>Specify a method to gather training data</td>
</tr>
<tr>
<td></td>
<td>dace_method_pointer</td>
<td></td>
</tr>
<tr>
<td></td>
<td>actual_model_pointer</td>
<td>A surrogate model pointer that guides a method to whether it should use a surrogate model or compute truth function evaluations</td>
</tr>
<tr>
<td>Optional</td>
<td>reuse_points</td>
<td>Surrogate model training data reuse control</td>
</tr>
<tr>
<td>Optional</td>
<td>import_build_points_file</td>
<td>File containing points you wish to use to build a surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>export_approx_points_file</td>
<td>Output file for surrogate model value evaluations</td>
</tr>
</tbody>
</table>
Description

The global surrogate model requires specification of one of the following approximation types:

1. Polynomial
2. Gaussian process (Kriging interpolation)
3. Layered perceptron artificial neural network approximation
4. MARS
5. Moving least squares
6. Radial basis function
7. Voronoi Piecewise Surrogate (VPS)

All these approximations are implemented in SurfPack[36], except for VPS. In addition, a second version of Gaussian process is implemented directly in Dakota.

Training Data

Training data can be taken from prior runs, stored in a datafile, or by running a Design of Experiments method. The keywords listed below are used to determine how to collect training data:

- `dace_method_pointer`
- `reuse_points`
- `import_points_file`
- `use_derivatives` The source of training data is determined by the contents of a provided `import_points_file`, whether `reuse_points` and `use_derivatives` are specified, and the contents of the method block specified by `dace_method_pointer`. `use_derivatives` is a special case, the other keywords are discussed below.

The number of training data points used in building a global approximation is determined by specifying one of three point counts:

1. `minimum_points`: minimum required or minimum "reasonable" amount of training data. Defaults to \(d+1\) for \(d\) input dimensions for most models, e.g., polynomials override to the number of coefficients required to estimate the requested order.
2. `recommended_points`: recommended number of training data, (this is the default option, if none of the keywords is specified). Defaults to \(5\times d\), except for polynomials where it's equal to the minimum.
3. **total_points**: specify the number of training data points. However, if the `total_points` value is less than the default `minimum_points` value, the `minimum_points` value is used.

The sources of training data depend on the number of training points, $N_t$, the number of points in the import file, $N_i$, and the value of `reuse_points`.

- If there is no import file, all training data come from the DACE method
- If there is an import file, all $N_i$ points from the file are used, and the remaining $N_t - N_i$ points come from the DACE method
- If there is an import file and `reuse_points` is:
  - `none` - all $N_t$ points from DACE method
  - `region` - only the points within a trust region are taken from the import file, and all remaining points are from the DACE method.
  - `all` - (Default) all $N_i$ points from the file are used, and the remaining $N_t - N_i$ points come from the DACE method

**Surrogate Correction**

A correction model can be added to the constructed surrogate in order to better match the training data. The specified correction method will be applied to the surrogate, and then the corrected surrogate model is used by the method.

Finally, the quality of the surrogate can be tested using the `metrics` and `challenge_points_file` keywords.

**Theory**

Global methods, also referred to as response surface methods, involve many points spread over the parameter ranges of interest. These surface fitting methods work in conjunction with the sampling methods and design of experiments methods.

**Procedures for Surface Fitting**

The surface fitting process consists of three steps:

1. selection of a set of design points
2. evaluation of the true response quantities (e.g., from a user-supplied simulation code) at these design points,
3. using the response data to solve for the unknown coefficients (e.g., polynomial coefficients, neural network weights, kriging correlation factors) in the surface fit model.

In cases where there is more than one response quantity (e.g., an objective function plus one or more constraints), then a separate surface is built for each response quantity. Currently, the surface fit models are built using only 0th-order information (function values only), although extensions to using higher-order information (gradients and Hessians) are possible.

Each surface fitting method employs a different numerical method for computing its internal coefficients. For example, the polynomial surface uses a least-squares approach that employs a singular value decomposition to compute the polynomial coefficients, whereas the kriging surface uses Maximum Likelihood Estimation to compute its correlation coefficients. More information on the numerical methods used in the surface fitting codes is provided in the Dakota Developers Manual.

**See Also**

These keywords may also be of interest:
• local
• hierarchical
• multipoint

7.3.3.2.1  experimental_gaussian_process

• Keywords Area
• model
• surrogate
• global
• experimental_gaussian_process

Use the Gaussian process regression surrogate from the surrogates module.

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>Trend</td>
<td>trend</td>
<td>This keyword enables the use of deterministic polynomial trend function</td>
</tr>
<tr>
<td>Optional</td>
<td>num_restarts</td>
<td></td>
<td>Number of optimization restarts for L-BFGS-B</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Nugget (Group 1)</td>
<td>nugget</td>
<td>Value for the fixed nugget parameter</td>
</tr>
<tr>
<td></td>
<td></td>
<td>find_nugget</td>
<td>Use regression to estimate the nugget.</td>
</tr>
<tr>
<td>Optional</td>
<td>options_file</td>
<td></td>
<td>Filename for a YAML file that specifies Gaussian process options</td>
</tr>
</tbody>
</table>
Description

This Gaussian process implementation is contained in Dakota's surrogates module and is considered experimental. It uses gradient-based optimization with restarts to determine hyperparameters and trend coefficients. Nugget and trend estimation are optional.

7.3.3.2.1.1 trend

- Keywords Area
  - model
  - surrogate
  - global
  - experimental_gaussian_process
  - trend

This keyword enables the use of deterministic polynomial trend function

Specification

Alias: none

Argument(s): none

Default: reduced_quadratic

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Trend Order (Group 1)</td>
<td>constant</td>
<td>Constant trend function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>linear</td>
<td>Use a linear polynomial or trend function</td>
</tr>
</tbody>
</table>
Description

Enable the use of a deterministic polynomial trend function whose coefficients are found through regression. The user may choose a constant, linear, or quadratic polynomial.

Default Behavior  Defaults to no trend when keyword is absent.

7.3.3.2.1  none

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- trend
- none

No trend function

Specification

Alias: none

Argument(s): none

Description

See parent page

7.3.3.2.1.2  constant

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- trend
- constant

Constant trend function
7.3 model

Specification

Alias: none
Argument(s): none

Description

See parent page

7.3.3.2.1.4 linear

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- trend
- linear

Use a linear polynomial or trend function

Specification

Alias: none
Argument(s): none

Description

See parent page

7.3.3.2.1.5 reduced_quadratic

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- trend
- reduced_quadratic

Quadratic polynomials - main effects only

Specification

Alias: none
Argument(s): none
Description

In 2 or more dimensions, this polynomial omits the interaction, or mixed, terms.

7.3.3.2.1.6 quadratic

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- trend
- quadratic

Use a quadratic polynomial or trend function

Specification

Alias: none
Argument(s): none

Description

See parent page

7.3.3.2.1.7 num_restarts

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- num_restarts

Number of optimization restarts for L-BFGS-B

Specification

Alias: none
Argument(s): INTEGER

Description

Number of optimization runs at random initial guesses for maximum likelihood estimation. Multiple restarts are used to increase the chance of finding the global minimum with a gradient-based (i.e. local) optimization method.

Default Behavior  Default value of 20 restarts.
7.3 model

7.3.3.2.1.8 nugget

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- nugget

Value for the fixed nugget parameter

Specification

Alias: none
Argument(s): REAL
Default: None

Description

This keyword is used to specify a value for the nugget parameter. Its presence will turn off nugget estimation.

Default Behavior
No nugget when this keyword is absent. Alternatively a nugget may be estimated as part of the regression procedure.

7.3.3.2.1.9 find_nugget

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- find_nugget

Use regression to estimate the nugget.

Specification

Alias: none
Argument(s): INTEGER
Default: None

Description

Estimate a nugget term in the Gaussian process Gram matrix.

Default Behavior
Off by default. The user may alternatively specify a fixed nugget.
7.3.3.2.1.10   options_file

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- options_file

Filename for a YAML file that specifies Gaussian process options

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** no advanced options file

**Description**

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the Gaussian process that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

**Examples**

```yaml
# FILE: gp_options.yaml

GP Parameters:
  Sigma Bounds:
    lower bound: 1.0e-2
    upper bound: 1.0e2
  scaler name: standardization
  num restarts: 20
Nugget:
  fixed nugget: 0.0
  estimate nugget: true
  Bounds:
    lower bound: 3.17e-8
    upper bound: 1.0e-2
gp seed: 42
standardize response: false
Trend:
  estimate trend: true
Options:
  max degree: 2
  reduced basis: false
  p-norm: 1.0
  scaler type: none
  regression solver type: SVD
  verbosity: 1
kernel type: squared exponential
Length-scale Bounds:
  lower bound: 1.0e-2
  upper bound: 1.0e2
verbosity: 1
```
7.3.3.2.1.11 export_approx_variance_file

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **experimental_gaussian_process**
- **export_approx_variance_file**

Output file for surrogate model variance evaluations

**Specification**

**Alias:** none  
**Argument(s):** STRING  
**Default:** no variance export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose One</strong></td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response variance values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**See Also**

These keywords may also be of interest:

- **export_approx_points_file**
7.3.3.2.1.12 custom_annotated

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_approx_variance_file
- custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.
Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 0.9 1.1 0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.3.3.2.1.13 header

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_approx_variance_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none
Argument(s): none

Description

See description of parent custom_annotated
7.3.3.2.1.14 eval_id

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_approx_variance_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent custom_annotated

7.3.3.2.1.15 interface_id

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_approx_variance_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent custom_annotated
7.3.3.2.1.16 annotated

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_approx_variance_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom.annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
    tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

```plaintext
%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.3.3.2.1.17 freeform

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **experimental_gaussian_process**
- **export_approx_variance_file**
- **freeform**

Selects freeform file format

Topics

This keyword is related to the topics:

- **file_formats**

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The **annotated** format is the default for tabular export/import. To change this behavior, specify **freeform** in the relevant export/import context.

**Usage Tips**
- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.3.3.2.1.18 export_model

- Keywords Area
  - model
  - surrogate
  - global
  - experimental_gaussian_process
  - export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the `formats` specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive** - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive** - Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file** - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console** - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Funtions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used.

Currently, no other surrogate models can be exported.

#### Default Behavior

No export.

#### Expected Output

Output depends on selected format; see the `formats` specification.

#### Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.1.19  `filename_prefix`
User-customizable portion of exported/imported surrogate model filenames

Topics
This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description
Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples
This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```
surrogate global gaussian_process surfpack
   export_model
      filename_prefix = 'my_surrogate'
      formats
         algebraic_file
         binary_archive
```

7.3.3.2.1.20 formats

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- export_model
Formats for surrogate model export

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted.

See export_model and the entries for the format selection keywords for further information.

7.3.3.2.1.21 text_archive

• Keywords Area
• model
• surrogate
• global
• experimental_gaussian_process
• export_model
• formats
• text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

• surrogate_models
7.3 model

Specification

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`.

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which `sps` stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

**Examples**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.1.22 binary_archive

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **experimental_gaussian_process**
- **export_model**
- **formats**
- **binary_archive**

Surrogate model binary archive file format

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named `{prefix}.{response_descriptor}.bin`.
Other (Surfpack) surrogates will export/import using files named \( \text{prefix}.\{\text{response\_descriptor}\}.bsps \), in which 'bsps' stands for binary Surfpack surrogate.

See \texttt{filename\_prefix} for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.1.23 \texttt{import\_model}

- \texttt{Keywords Area}
- \texttt{model}
- \texttt{surrogate}
- \texttt{global}
- \texttt{experimental\_gaussian\_process}
- \texttt{import\_model}

Import surrogate model from archive file

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: none

\textbf{Child Keywords}:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>\texttt{filename_prefix}</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
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</table>

\textbf{Required (Choose One)}

<table>
<thead>
<tr>
<th>Surrogate Import Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>\texttt{text_archive}</td>
<td></td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>\texttt{binary_archive}</td>
<td></td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Imports a surrogate model from a file in \texttt{binary\_archive} or \texttt{text\_archive} format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords \texttt{filename\_prefix} and \texttt{binary\_archive} or \texttt{text\_archive}.

\textbf{Default Behavior}
When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through `dace_method_pointer`.

Usage Tips

When importing a surrogate model, it is crucial that the `global` surrogate model part of the Dakota input file be identical for export and import, except for changing `export_model` and its child keywords to those needed for `import_model`. Any other keywords such as specifying a `dace_iterate` or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed `initial_state` values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the `surfpack` binary or the experimental `dakota.surrogates` Python module, it is untested and not the primary use case for this capability.

Examples

The following model block exports a Gaussian process after it’s built:

```dakota
model
  id_model = 'SURR_M'
  surrogate global
    dace_method_pointer = 'DACE'
    experimental_gaussian_process
      metrics 'root_mean_squared'
      cross_validation folds 5
      export_model
        filename_prefix = 'gp_export'
        formats = binary_archive
        export_approx_points_file 'gp_values.dat'
        export_approx_variance_file 'gp_variance.dat'

And this model block imports it using a nearly identical model specification:

```dakota
model
  id_model = 'SURR_M'
  surrogate global
    dace_method_pointer = 'DACE'
    experimental_gaussian_process
      metrics 'root_mean_squared'
      cross_validation folds 5
    import_model
      filename_prefix = 'gp_export'
      binary_archive
      export_approx_points_file 'gp_values.dat'
      export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.1.24  filename_prefix

- Keywords Area
- model
- surrogate
User-customizable portion of exported/imported surrogate model filenames

Topics
This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description
Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples
This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```python
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
  formats
    algebraic_file
    binary_archive
```

7.3.3.2.1.25  text_archive

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- import_model
Surrogate model plain-text archive file format

Topics
This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description
When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples
The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.1.26 binary_archive

- Keywords Area
- model
- surrogate
- global
- experimental_gaussian_process
- import_model
- binary_archive

Surrogate model binary archive file format

Topics
This keyword is related to the topics:

- surrogate_models
Specification

**Alias:** none

**Argument(s):** none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.2 gaussian_process

- **Keywords Area**

- **model**

- **surrogate**

- **global**

- **gaussian_process**

Gaussian Process surrogate model

Specification

**Alias:** kriging

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
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<td>dakota</td>
<td>Select the built in Gaussian Process surrogate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>surfpack</td>
<td>Use the Surfpack version of Gaussian Process surrogates</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>export_approx_variance_file</td>
<td>Output file for surrogate model variance evaluations</td>
</tr>
</tbody>
</table>
Description

Use the Gaussian process (GP) surrogate from Surfpack, which is specified using the surfpack keyword. An alternate version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

### 7.3.3.2.2.1 dakota

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **gaussian_process**
- **dakota**

Select the built in Gaussian Process surrogate

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>point_selection</td>
<td></td>
<td>Enable greedy selection of well-spaced build points</td>
</tr>
<tr>
<td>Optional</td>
<td>trend</td>
<td></td>
<td>Choose a trend function for a Gaussian process surrogate</td>
</tr>
</tbody>
</table>

**Description**

A second version of GP surrogates was available in prior versions of Dakota. **For now, both versions are supported but the dakota version is deprecated and intended to be removed in a future release.**

Historically these models were drastically different, but in Dakota 5.1, they became quite similar. They now differ in that the Surfpack GP has a richer set of features/options and tends to be more accurate than the Dakota version. Due to how the Surfpack GP handles ill-conditioned correlation matrices (which significantly contributes to its greater accuracy), the Surfpack GP can be a factor of two or three slower than Dakota’s. As of Dakota 5.2, the Surfpack implementation is the default in all contexts except Bayesian calibration.

More details on the gaussian_process dakota model can be found in[60].
Dakota’s GP deals with ill-conditioning in two ways. First, when it encounters a non-invertible correlation matrix it iteratively increases the size of a “nugget,” but in such cases the resulting approximation smooths rather than interpolates the data. Second, it has a `point_selection` option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

This differs from the `point_selection` option of the Dakota GP which initially chooses a well-spaced subset of points and finds the correlation parameters that are most likely for that one subset.

### 7.3.3.2.2.2 point_selection

Enable greedy selection of well-spaced build points

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

*Alias*: none

*Argument(s)*: none

*Default*: no point selection

**Description**

The Dakota Gaussian Process model has a `point_selection` option (default off) that uses a greedy algorithm to select a well-spaced subset of points prior to the construction of the GP. In this case, the GP will only interpolate the selected subset. Typically, one should not need point selection in trust-region methods because a small number of points are used to develop a surrogate within each trust region. Point selection is most beneficial when constructing with a large number of points, typically more than order one hundred, though this depends on the number of variables and spacing of the sample points.

### 7.3.3.2.2.3 trend

- Keywords Area
- model
- surrogate
Choose a trend function for a Gaussian process surrogate

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** reduced_quadratic

**Child Keywords:**

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required (Choose One)</td>
<td>Trend Order (Group 1)</td>
<td>constant</td>
<td>Constant trend function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>linear</td>
<td>Use a linear polynomial or trend function</td>
</tr>
<tr>
<td></td>
<td></td>
<td>reduced_quadratic</td>
<td>Quadratic polynomials - main effects only</td>
</tr>
</tbody>
</table>

**Description**

The only trend functions that are currently supported are polynomials.

The trend function is selected using the `trend` keyword, with options `constant`, `linear`, or `reduced_quadratic`. The `reduced_quadratic` trend function includes the main effects, but not mixed/interaction terms. The Surfpack GP (See surfpack) has the additional option of (a full) `quadratic`.

7.3.3.2.2.4 constant

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- dakota
- trend
- constant

Constant trend function
Use a linear polynomial or trend function

Quadratic polynomials - main effects only
7.3 model

Specification

Alias: none
Argument(s): none

Description

In 2 or more dimensions, this polynomial omits the interaction, or mixed, terms.

7.3.3.2.2.7 surfpack

- Keywords
  - Area
  - model
  - surrogate
  - global
  - gaussian_process
  - surfpack

Use the Surfpack version of Gaussian Process surrogates

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>trend</td>
<td>trend</td>
<td>Choose a trend function for a Gaussian process surrogate</td>
</tr>
<tr>
<td>Optional</td>
<td>optimization_method</td>
<td>optimization_method</td>
<td>Change the optimization method used to compute hyperparameters</td>
</tr>
<tr>
<td>Optional</td>
<td>max_trials</td>
<td>max_trials</td>
<td>Max number of likelihood function evaluations</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>nugget (Group 1)</td>
<td>nugget</td>
<td>Specify a nugget to handle ill-conditioning</td>
</tr>
<tr>
<td></td>
<td>find_nugget</td>
<td></td>
<td>Have Surfpack compute a nugget to handle ill-conditioning</td>
</tr>
</tbody>
</table>
This keyword specifies the use of the Gaussian process that is incorporated in our surface fitting library called Surfpack.

Several user options are available:

1. Optimization methods:
   Maximum Likelihood Estimation (MLE) is used to find the optimal values of the hyper-parameters governing the trend and correlation functions. By default the global optimization method DIRECT is used for MLE, but other options for the optimization method are available. See optimization_method.
   The total number of evaluations of the likelihood function can be controlled using the max_trials keyword followed by a positive integer. Note that the likelihood function does not require running the "truth" model, and is relatively inexpensive to compute.

2. Trend Function:
   The GP models incorporate a parametric trend function whose purpose is to capture large-scale variations. See trend.

3. Correlation Lengths:
   Correlation lengths are usually optimized by Surfpack, however, the user can specify the lengths manually. See correlation_lengths.

4. Ill-conditioning
   One of the major problems in determining the governing values for a Gaussian process or Kriging model is the fact that the correlation matrix can easily become ill-conditioned when there are too many input points close together. Since the predictions from the Gaussian process model involve inverting the correlation matrix, ill-conditioning can lead to poor predictive capability and should be avoided.
   Note that a sufficiently bad sample design could require correlation lengths to be so short that any interpolatory GP model would become inept at extrapolation and interpolation.
   The surfpack model handles ill-conditioning internally by default, but behavior can be modified using:

5. Gradient Enhanced Kriging (GEK).
   The use_derivatives keyword will cause the Surfpack GP to be constructed from a combination of function value and gradient information (if available).
   See notes in the Theory section.

Theory

**Gradient Enhanced Kriging**

Incorporating gradient information will only be beneficial if accurate and inexpensive derivative information is available, and the derivatives are not infinite or nearly so. Here "inexpensive" means that the cost of evaluating a
function value plus gradient is comparable to the cost of evaluating only the function value, for example gradients computed by analytical, automatic differentiation, or continuous adjoint techniques. It is not cost effective to use derivatives computed by finite differences. In tests, GEK models built from finite difference derivatives were also significantly less accurate than those built from analytical derivatives. Note that GEK's correlation matrix tends to have a significantly worse condition number than Kriging for the same sample design.

This issue was addressed by using a pivoted Cholesky factorization of Kriging's correlation matrix (which is a small sub-matrix within GEK's correlation matrix) to rank points by how much unique information they contain. This reordering is then applied to whole points (the function value at a point immediately followed by gradient information at the same point) in GEK's correlation matrix. A standard non-pivoted Cholesky is then applied to the reordered GEK correlation matrix and a bisection search is used to find the last equation that meets the constraint on the (estimate of) condition number. The cost of performing pivoted Cholesky on Kriging's correlation matrix is usually negligible compared to the cost of the non-pivoted Cholesky factorization of GEK's correlation matrix. In tests, it also resulted in more accurate GEK models than when pivoted Cholesky or whole-point-block pivoted Cholesky was performed on GEK's correlation matrix.

7.3.3.2.2.8 trend

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- trend

Choose a trend function for a Gaussian process surrogate

Specification

Alias: none

Argument(s): none

Default: reduced_quadratic

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
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<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td>Trend Order (Group 1)</td>
<td>constant</td>
<td>Constant trend function</td>
</tr>
<tr>
<td>linear</td>
<td>Use a linear polynomial or trend function</td>
<td></td>
<td></td>
</tr>
<tr>
<td>reduced_quadratic</td>
<td>Quadratic polynomials - main effects only</td>
<td></td>
<td></td>
</tr>
<tr>
<td>quadratic</td>
<td>Use a quadratic polynomial or trend function</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
The only trend functions that are currently supported are polynomials.

The trend function is selected using the trend keyword, with options constant, linear, or reduced_quadratic. The reduced_quadratic trend function includes the main effects, but not mixed/interaction terms. The Surfpack GP (See surfpack) has the additional option of (a full) quadratic.

### 7.3.3.2.9 constant

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **gaussian_process**
- **surfpack**
- **trend**
- **constant**

#### Constant trend function

**Description**

See parent page

#### Specification

**Alias**: none

**Argument(s)**: none

### 7.3.3.2.10 linear

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **gaussian_process**
- **surfpack**
- **trend**
- **linear**

#### Use a linear polynomial or trend function

**Specification**

**Alias**: none

**Argument(s)**: none
Description

See parent page

7.3.3.2.2.11 reduced_quadratic

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- trend
- reduced_quadratic

Quadratic polynomials - main effects only

Specification

**Alias:** none

**Argument(s):** none

Description

In 2 or more dimensions, this polynomial omits the interaction, or mixed, terms.

7.3.3.2.2.12 quadratic

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- trend
- quadratic

Use a quadratic polynomial or trend function

Specification

**Alias:** none

**Argument(s):** none
Description

See parent page

7.3.3.2.2.13 optimization_method

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- optimization_method

Change the optimization method used to compute hyperparameters

Specification

Alias: none

Argument(s): STRING

Default: global

Description

Select the optimization method to compute hyperparameters of the Gaussian Process by specifying one of these arguments:

- global (default) - DIRECT method
- local - CONMIN method
- sampling - generates several random guesses and picks the candidate with greatest likelihood
- none - no optimization, pick the center of the feasible region

The none option, and the starting location of the local optimization, default to the center, in log(correlation length) scale, of the feasible region.

Surfpack picks a small feasible region of correlation parameters.

Note that we have found the global optimization method to be the most robust.

7.3.3.2.2.14 max_trials

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- max_trials

Max number of likelihood function evaluations
Specify a nugget to handle ill-conditioning

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**  
See parent page

7.3.3.2.2.15 nugget

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- nugget

Specify a nugget to handle ill-conditioning

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** None

**Description**  
By default, the Surfpack GP handles ill-conditioning and does not use a nugget. If the user wishes to specify a nugget, there are two approaches.

- The user can specify the value of a nugget with **nugget**.
- Have Surfpack find the optimal value of the nugget. This is specified by **find_nugget**. There are two options for **find_nugget**.
  - **find_nugget = 1**: assume that the reciprocal condition number of the correlation matrix R, rcondR, is zero and calculate the nugget needed to make the worst case of R not ill-conditioned.
  - **find_nugget = 2**: calculate rcondR, which requires a Cholesky factorization. If rcondR indicates that R is not ill-conditioned, then kriging uses the Cholesky factorization. Otherwise, if rcondR says R is ill conditioned, then kriging will calculate the nugget needed to make the worst case of R not ill conditioned.

**find_nugget = 1** and **2** are similar, the second option just takes more computation (the initial Cholesky factorization) for larger problems.
8.3.3.2.2.16 find_nugget

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- find_nugget

Have Surfpack compute a nugget to handle ill-conditioning

Specification

Alias: none

Argument(s): INTEGER

Default: None

Description

By default, the Surfpack GP handles ill-conditioning and does not use a nugget. If the user wishes to specify a nugget, there are two approaches.

- The user can specify the value of a nugget with nugget.
- Have Surfpack find the optimal value of the nugget. This is specified by find_nugget. There are two options for find_nugget.
  - find_nugget = 1: assume that the reciprocal condition number of the correlation matrix R, rcondR, is zero and calculate the nugget needed to make the worst case of R not ill-conditioned.
  - find_nugget = 2: calculate rcondR, which requires a Cholesky factorization. If rcondR indicates that R is not ill-conditioned, then kriging uses the Cholesky factorization. Otherwise, if rcondR says R is ill conditioned, then kriging will calculate the nugget needed to make the worst case of R not ill conditioned.

find_nugget = 1 and 2 are similar, the second option just takes more computation (the initial Cholesky factorization) for larger problems.

7.3.3.2.2.17 correlation_lengths

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- correlation_lengths

Specify the correlation lengths for the Gaussian process
7.3 model

Specification

Alias: none
Argument(s): REALLIST
Default: internally computed correlation_lengths

Description

Directly specify \texttt{correlation\_lengths} as a list of \texttt{N} real numbers where \texttt{N} is the number of input dimensions.

7.3.3.2.2.18 export_model

- Keywords Area
- model
- surrogate
- global
- \texttt{gaussian\_process}
- surfpack
- export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- \texttt{surrogate\_models}

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>\texttt{filename_prefix}</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>\texttt{formats}</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>
Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- text_archive - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- binary_archive - Binary, machine-readable archive for re-import or use with the surfpack executable
- algebraic_file - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- algebraic_console - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword gaussian_process surfpack)
- Artificial neural network (keyword neural_network)
- Radial basis Functions (keyword radial_basis)
- Polynomial (keyword polynomial)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword mars) and moving least squares (keyword moving_least_squares) models, only text_archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior

No export.

Expected Output

Output depends on selected format; see the formats specification.

Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias**: none

**Argument(s)**: STRING

**Default**: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.extension`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
  formats
    algebraic_file
    binary_archive
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
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<th>Dakota Keyword Description</th>
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<td>Optional</td>
<td>text_archive</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_file</td>
<td>algebraic_file</td>
<td>Export surrogate model in algebraic format to a file</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_console</td>
<td>algebraic_console</td>
<td>Export surrogate model in algebraic format to the console</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted.

See export_model and the entries for the format selection keywords for further information.

7.3.3.2.21 text_archive

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format
7.3 model

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.22 binary_archive

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none
Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.
Experimental surrogates will export to (import from) binary archives named {prefix}.{response_descriptor}.bin.
Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.bsps, in which ‘bsps’ stands for binary Surfpack surrogate.
See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.23 algebraic_file

Export surrogate model in algebraic format to a file

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to a file in a human-readable “algebraic” format. The file is named using the pattern {prefix}.{response_descriptor}.alg. See filename_prefix for further information about exported surrogate file naming. The file contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.
The output written to the file matches exactly the output written to the console when `algebraic_console` is specified.

7.3.3.2.2.24  algebraic_console

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- export_model
- formats
- algebraic_console

Export surrogate model in algebraic format to the console

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none

**Argument(s):** none

**Description**

After the surrogate model has been built, Dakota will export it to the console (screen, or output file if Dakota was run using the `-o` option) in a human-readable “algebraic” format. The output contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

**Expected Output**

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the screen for the exported model matches exactly the output written to file when `algebraic_file` is specified. Use of `algebraic_file` is preferred over `algebraic_console`, which exists largely to provide a measure of backward compatibility.

7.3.3.2.2.25  import_model

- Keywords Area
- model
- surrogate
- global
Import surrogate model from archive file

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td><strong>Optional</strong></td>
<td></td>
<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
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<td><strong>Required (Choose One)</strong></td>
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<td>text_archive</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

**Description**

Imports a surrogate model from a file in `binary_archive` or `text_archive` format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords `filename_prefix` and `binary_archive` or `text_archive`.

**Default Behavior**

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through `dace_method_pointer`.

**Usage Tips**

When importing a surrogate model, it is crucial that the `global` surrogate model part of the Dakota input file be identical for export and import, except for changing `export_model` and its child keywords to those needed for `import_model`. Any other keywords such as specifying a `dace_iterator` or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed `initial_state` values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped...
when evaluating it. So it is important the state values for s and t haven't changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the `surfpack` binary or the experimental `dakota.surrogates` Python module, it is untested and not the primary use case for this capability.

Examples

The following model block exports a Gaussian process after it's built:

```plaintext
model
id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
metrics 'root_mean_squared'
cross_validation folds 5
export_model
filename_prefix = 'gp_export'
formats = binary_archive
export_approx_points_file 'gp_values.dat'
export_approx_variance_file 'gp_variance.dat'
```

And this model block imports it using a nearly identical model specification:

```plaintext
model
id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
metrics 'root_mean_squared'
cross_validation folds 5
import_model
filename_prefix = 'gp_export'
formats = binary_archive
export_approx_points_file 'gp_values.dat'
export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.2.26 `filename_prefix`

- Keywords Area
- `model`
- `surrogate`
- `global`
- `gaussian_process`
- `surfpack`
- `import_model`
- `filename_prefix`

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- `surrogate_models`
Keywords Area

Specification

**Alias:** none  

**Argument(s):** STRING  

**Default:** exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern. The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```
surrogate global gaussian_process surfpack  
   export_model  
      filename_prefix = ’my_surrogate’  
      formats  
         algebraic_file  
         binary_archive
```

7.3.3.2.2.27 text_archive

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none  

**Argument(s):** none
Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt.` Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.28 binary_archive

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- surfpack
- import_model
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named `{prefix}.{response_descriptor}.bin.` Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which ‘bsps’ stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
7.3.3.2.29 export_approx_variance_file

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- export_approx_variance_file

Output file for surrogate model variance evaluations

**Specification**

Alias: none

Argument(s): STRING

Default: no variance export to a file

Child Keywords:

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
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<td>Selects annotated tabular file format</td>
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<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response variance values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

**Usage Tips**

Dakota exports tabular data in one of three formats:

- annotated (default)
- custom_annotated
- freeform

**See Also**

These keywords may also be of interest:

- export_approx_points_file

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7.3.3.2.30 custom_annotated

- **Keywords Area**
- model
- surrogate
- global
- gaussian_process
- export_approx_variance_file
- custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- **file_formats**

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
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<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
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<td>header</td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.
Usage Tips

• Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id   x1   x2   obj_fn   nln_ineq_con_1   nln_ineq_con_2
1   0.9   1.1   0.0002   0.26   0.76
2   0.90009   1.1   0.0001996404857   0.2601620081   0.759955
3   0.89991   1.1   0.0002003604863   0.2598380081   0.760045
...```

7.3.2.2.31 header

• Keywords Area

• model

• surrogate

• global

• gaussian_process

• export_approx_variance_file

• custom_annotated

• header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated
7.3.2.2.32 eval_id

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- export_approx_variance_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

See description of parent custom_annotated

7.3.2.2.33 interface_id

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- export_approx_variance_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

See description of parent custom_annotated
7.3.3.2.2.34  annotated

- Keywords Area
- model
- surrogate
- global
- gaussian_process
- export_approx_variance_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The **annotated** keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify **custom.annotated header eval_id**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to **annotated format**, though **freeform** remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the **freeform** option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    annotated
```

Resulting tabular file:

```plaintext
%eval_id interface  x1  x2  obj_fn  nln_ineq_con_1  nln_ineq_con_2
1   NO_ID     0.9  1.1  0.0002  0.26     0.76
2   NO_ID     0.90009 1.1  0.0001996404857 0.2601620081 0.759955
3   NO_ID     0.89991 1.1  0.0002003604863 0.2598380081 0.760045
...```

7.3.3.2.2.35 freeform

• Keywords Area

• model

• surrogate

• global

• gaussian_process

• export_approx_variance_file

• freeform

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips
Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
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<td>0.0002003604863</td>
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</tr>
<tr>
<td></td>
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</tr>
</tbody>
</table>

7.3.3.2.3 mars

Keywords Area

model

surrogate

global

mars

Multivariate Adaptive Regression Spline (MARS)

Specification

Alias: none

Argument(s): none

Child Keywords:
### Description

This surface fitting method uses multivariate adaptive regression splines from the MARS3.5 package[27] developed at Stanford University.

The MARS reference material does not indicate the minimum number of data points that are needed to create a MARS surface model. However, in practice it has been found that at least $n_{quad}$, and sometimes as many as 2 to 4 times $n_{quad}$, data points are needed to keep the MARS software from terminating. Provided that sufficient data samples can be obtained, MARS surface models can be useful in SBO and OUU applications, as well as in the prediction of global trends throughout the parameter space.

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

### Theory

The form of the MARS model is based on the following expression:

$$\hat{f}(x) = \sum_{m=1}^{M} a_m B_m(x)$$

where the $a_m$ are the coefficients of the truncated power basis functions $B_m$, and $M$ is the number of basis functions. The MARS software partitions the parameter space into subregions, and then applies forward and backward regression methods to create a local surface model in each subregion. The result is that each subregion contains its own basis functions and coefficients, and the subregions are joined together to produce a smooth, $C^2$-continuous surface model.

MARS is a nonparametric surface fitting method and can represent complex multimodal data trends. The regression component of MARS generates a surface model that is not guaranteed to pass through all of the response data values. Thus, like the quadratic polynomial model, it provides some smoothing of the data.

#### 7.3.3.2.3.1 max_bases

- Keywords Area
- model
- surrogate
• global
• mars
• max_bases

Maximum number of MARS bases

Specification

Alias: none
Argument(s): INTEGER

Description

The maximum number of basis functions allowed in the MARS approximation model.

7.3.3.2.3.2 Interpolation

• Keywords Area
• model
• surrogate
• global
• mars
• interpolation

MARS model interpolation type

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Interpolation Order (Group 1)</th>
<th>Dakota Keyword</th>
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<td></td>
<td>linear</td>
<td>Linear interpolation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>cubic</td>
<td>Cubic interpolation</td>
</tr>
</tbody>
</table>

Description

The MARS model interpolation type: linear or cubic.

7.3.3.2.3.3 linear

• Keywords Area
• model
• surrogate
Linear interpolation

Specification

**Alias:** none

**Argument(s):** none

Description

Use linear interpolation in the MARS model.

7.3.3.2.3.4 cubic

**Keywords** Area

**model**

**surrogate**

**global**

**mars**

**interpolation**

**cubic**

Cubic interpolation

Specification

**Alias:** none

**Argument(s):** none

Description

Use cubic interpolation in the MARS model.

7.3.3.2.3.5 export_model

**Keywords** Area

**model**

**surrogate**

**global**

**mars**

**export_model**

Exports surrogate model in user-specified format(s)
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Required</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
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<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>formats</td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- text_archive: Plain-text, machine-readable archive for re-import or use with the surfpack executable
- binary_archive: Binary, machine-readable archive for re-import or use with the surfpack executable
- algebraic_file: Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- algebraic_console: Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword gaussian_process surfpack)
- Artificial neural network (keyword neural_network)
- Radial basis Funtions (keyword radial_basis)
- Polynomial (keyword polynomial)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword mars) and moving least squares (keyword moving_least_squares) models, only text_archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior
No export.

**Expected Output**

Output depends on selected format; see the *formats* specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

### 7.3.3.2.3.6 filename_prefix

- **Keywords Area**
  - model
  - surrogate
  - global
  - mars
  - export_model
  - filename_prefix

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** exported_surrogate

**Description**

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response *descriptors* provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

**Examples**

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive

7.3.3.2.3.7 formats
   • Keywords Area
   • model
   • surrogate
   • global
   • mars
   • export_model
   • formats

Formats for surrogate model export

Topics

This keyword is related to the topics:
   • surrogate_models

Specification

Alias: none
Argument(s): none
Child Keywords:

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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>binary archive file format</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted.
See export_model and the entries for the format selection keywords for further information.
Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export (import from) text archives named `{prefix}.{response_descriptor}.txt`.

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
• formats
  • binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:
  • surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.bsps, in which ‘bsps’ stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.3.10  import_model
  • Keywords Area
  • model
  • surrogate
  • global
  • mars
  • import_model

Import surrogate model from archive file

Specification

Alias: none
Argument(s): none
Child Keywords:
<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tr>
<td></td>
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<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

**Description**

Imports a surrogate model from a file in binary_archive or text_archive format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords filename_prefix and binary_archive or text_archive.

**Default Behavior**

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through dace_method_pointer.

**Usage Tips**

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing export_model and its child keywords to those needed for import_model. Any other keywords such as specifying a dace_iterator or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed initial_state values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the surfpack binary or the experimental_dakota.surrogates Python module, it is untested and not the primary use case for this capability.

**Examples**

The following model block exports a Gaussian process after it’s built:

```plaintext
model
    id_model = ‘SURR_M’
    surrogate global
        dace_method_pointer = ‘DACE’
        experimental_gaussian_process
```
And this model block imports it using a nearly identical model specification:

```plaintext
model
  id_model = 'SURR_M'
  surrogate global
  dace_method_pointer = 'DACE'
  experimental_gaussian_process
  metrics 'root_mean_squared'
  cross_validation folds 5
import_model
  filename_prefix = 'gp_export'
  binary_archive
  export_approx_points_file 'gp_values.dat'
  export_approx_variance_file 'gp_variance.dat'
```

### 7.3.3.2.3.11 filename_prefix

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **mars**
- **import_model**
- **filename_prefix**

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** exported_surrogate

**Description**

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.
Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
forms
  algebraic_file
  binary_archive
```

7.3.3.2.3.12 text_archive

- Keywords Area
- model
- surrogate
- global
- mars
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
7.3.3.2.3.13  binary_archive

- Keywords Area
- model
- surrogate
- global
- mars
- import_model
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named `{prefix}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which ‘bsps’ stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.4  moving_least_squares

- Keywords Area
- model
- surrogate
- global
- moving_least_squares

Moving Least Squares surrogate models
Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword</th>
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<td>Polynomial order for the MLS bases</td>
</tr>
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<td>Optional</td>
<td>weight_function</td>
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<td>Selects the weight function for the MLS model</td>
</tr>
<tr>
<td>Optional</td>
<td>export_model</td>
<td></td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
<tr>
<td>Optional</td>
<td>import_model</td>
<td></td>
<td>Import surrogate model from archive file</td>
</tr>
</tbody>
</table>

**Description**

Moving least squares is a further generalization of weighted least squares where the weighting is "moved" or recalculated for every new point where a prediction is desired[66].

**The implementation of moving least squares is still under development.** It tends to work well in trust region optimization methods where the surrogate model is constructed in a constrained region over a few points. The present implementation may not work as well globally.

Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

**Theory**

Moving Least Squares can be considered a more specialized version of linear regression models. In linear regression, one usually attempts to minimize the sum of the squared residuals, where the residual is defined as the difference between the surrogate model and the true model at a fixed number of points.

In weighted least squares, the residual terms are weighted so the determination of the optimal coefficients governing the polynomial regression function, denoted by \( \hat{f}(x) \), are obtained by minimizing the weighted sum of squares at \( N \) data points:

$$\sum_{n=1}^{N} w_n (|| \hat{f}(x_n) - f(x_n) ||)$$

7.3.3.2.4.1 basis_order

- Keywords Area
Polynomial order for the MLS bases

**Specification**

**Alias:** poly\_order  
**Argument(s):** INTEGER

**Description**

The polynomial order for the moving least squares basis function (default = 2).

7.3.3.2.4.2 weight\_function

- Keywords Area
- model
- surrogate
- global
- moving\_least\_squares
- weight\_function

Selects the weight function for the MLS model

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

The weight function decays as a function of distance from the training data. Specify one of:

- 1 (default): exponential decay in weight function; once differentiable MLS model
- 2: twice differentiable MLS model
- 3: three times differentiable MLS model
7.3.3.2.4.3 export_model

- **Keywords Area**
  - model
  - surrogate
  - global
  - moving_least_squares
  - export_model

Exports surrogate model in user-specified format(s)

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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</tr>
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<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

**Description**

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the `formats` specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive** - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive** - Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file** - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console** - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable
These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only `text_archive` and `binary_archive` formats may be used. Currently, no other surrogate models can be exported.

**Default Behavior**

No export.

**Expected Output**

Output depends on selected format; see the `formats` specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.4.4 `filename_prefix`

- **Keywords Area**
- `model`
- `surrogate`
- `global`
- `moving_least_squares`
- `export_model`
- `filename_prefix`

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

- `surrogate_models`

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
  export_model
    filename_prefix = 'my_surrogate'
    formats
      algebraic_file
      binary_archive
```

7.3.3.2.4.5 formats

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **moving_least_squares**
- **export_model**
- **formats**

Formats for surrogate model export

Topics

This keyword is related to the topics:

- **surrogate_models**

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See `export_model` and the entries for the format selection keywords for further information.

7.3.3.2.4.6 `text_archive`

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **moving_least_squares**
- **export_model**
- **formats**
- **text_archive**

Surrogate model plain-text archive file format

### Topics

This keyword is related to the topics:

- **surrogate_models**

### Specification

**Alias:** none

**Argument(s):** none

### Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt.`

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.
Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.4.7 binary_archive

- Keywords Area
- model
- surrogate
- global
- moving_least_squares
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none

**Argument(s):** none

**Description**

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named `{prefix}.{responseDescriptor}.bin`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{responseDescriptor}.bsps`, in which 'bsps' stands for binary Surfpack surrogate.

See [filename_prefix](#) for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.4.8 import_model

- Keywords Area
- model
- surrogate
- global
Import surrogate model from archive file

**Specification**

**Alias:** none  
**Argument(s):** none  
**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Surrogate Import Format (Group 1)</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

**Description**

Imports a surrogate model from a file in `binary_archive` or `text_archive` format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords `filename_prefix` and `binary_archive` or `text_archive`.

**Default Behavior**

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through `dace_method_pointer`.

**Usage Tips**

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing `export_model` and its child keywords to those needed for `import_model`. Any other keywords such as specifying a `dace_iterator` or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed `initial_state` values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.
While it might be possible to import surrogates externally-generated using the `surfpack` binary or the experimental `dakota.surrogates` Python module, it is untested and not the primary use case for this capability.

Examples

The following model block exports a Gaussian process after it's built:

```plaintext
model
id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
metrics 'root_mean_squared'
cross_validation folds 5
export_model
filename_prefix = 'gp_export'
formats = binary_archive
export_approx_points_file 'gp_values.dat'
export_approx_variance_file 'gp_variance.dat'
```

And this model block imports it using a nearly identical model specification:

```plaintext
model
id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
metrics 'root_mean_squared'
cross_validation folds 5
import_model
filename_prefix = 'gp_export'
binary_archive
export_approx_points_file 'gp_values.dat'
export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.4.9 filename_prefix

- Keywords Area
- model
- surrogate
- global
- moving_least_squares
- import_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models
Specification

Alias: none
Argument(s): STRING
Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
   export_model
      filename_prefix = 'my_surrogate'
      formats
         algebraic_file
         binary_archive
```

7.3.3.2.4.10 text_archive

- Keywords Area
- model
- surrogate
- global
- moving_least_squares
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none
7.3.3.2.4.11 binary_archive

- Keywords Area
- model
- surrogate
- global
- moving_least_squares
- import_model
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.
Experimental surrogates will export to (import from) binary archives named {prefix}.{response_descriptor}.bin.
Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.bsps, in which 'bsps' stands for Surfpack surrogate.
See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Global surrogate model based on functional tensor train decomposition

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>regression_type</td>
<td>Type of solver for forming function train approximations by regression</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_solver_-_iterations</td>
<td>Maximum iterations in determining polynomial coefficients</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>max_cross_-_iterations</td>
<td>Maximum number of iterations for cross-approximation during a rank adaptation.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>solver_tolerance</td>
<td>Convergence tolerance for the optimizer used during the regression solve.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>response_scaling</td>
<td>Perform bounds-scaling on response values prior to surrogate emulation</td>
</tr>
<tr>
<td>Optional</td>
<td>tensor_grid</td>
<td>Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.</td>
<td></td>
</tr>
<tr>
<td>-------------------</td>
<td>------------------------</td>
<td>------------------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>rounding_tolerance</td>
<td>An accuracy tolerance that is used to guide rounding during rank adaptation.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>arithmetic_- tolerance</td>
<td>A secondary rounding tolerance used for post-processing.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>start_order</td>
<td>(Initial) polynomial order of each univariate function within the functional tensor train.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>adapt_order</td>
<td>Activate adaptive procedure for determining the best basis order.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>kick_order</td>
<td>Increment used when adapting the basis order in function train methods.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_order</td>
<td>Maximum polynomial order of each univariate function within the functional tensor train.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>max_cv_order_candidates</td>
<td>Limit the number of cross-validation candidates for basis order.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>start_rank</td>
<td>The initial rank used for the starting point during a rank adaptation.</td>
<td></td>
</tr>
</tbody>
</table>
Description

Tensor train decompositions are approximations that exploit low rank structure in an input-output mapping. The form of the approximation can be written as a set of matrix valued products:

\[ f_r(x) = F_1(x_1) F_2(x_2) \ldots F_d(x_d) \]

Where the "cores" expand to

\[ F_k(x_k) = \begin{bmatrix} f_{11}^k(x_k) & \cdots & f_{r_k}^k(x_k) \\ \vdots & \ddots & \vdots \\ f_{r_k-1}^k(x_k) & \cdots & f_{r_k-r_k}^k(x_k) \end{bmatrix} \]

An example expansion over four random variables with rank vector \((1,7,5,3,1)\) is

\[ f_r(x) = \begin{bmatrix} f_{11}^{11}(x_1) & \cdots & f_{17}^{17}(x_1) \\ f_{21}^{11}(x_2) & \cdots & f_{25}^{15}(x_2) \\ \vdots & \ddots & \vdots \\ f_{31}^{11}(x_3) & \cdots & f_{33}^{13}(x_3) \\ \vdots & \ddots & \vdots \\ f_{41}^{11}(x_4) \end{bmatrix} \]

In the current implementation, orthogonal polynomial basis functions (Hermite and Legendre) are employed as the basis functions \(f_{jk}^{ij}(x_i)\), although the C3 library will enable additional options in the future.

The number of coefficients that must be computed by the regression solver can be inferred from the construction above. For each QoI, the regression size can be determined as follows:

- For \(v\) variables, orders \(o\) is a \(v\)-vector and ranks \(r\) is a \(v+1\)-vector
- the first core is a \(1 \times r_1\) row vector and contributes \((o_1+1)r_1\) terms
- the last core is a \(r_{v-1} \times 1\) col vector and contributes \((o_{v-1}+1)r_{v-1}\) terms
- the middle \(v-2\) cores are \(r_i \times r_{i+1}\) matrices that contribute \(r_i r_{i+1} (o_i+1)\) terms, \(i = 1, \ldots, v-2\)
- neighboring vec/mat dimensions must match, so there are \(v-1\) unique ranks

Usage Tips

This new capability is stabilizing and beginning to be embedded in higher-level strategies such as multilevel-multifidelity algorithms. It is not included in the Dakota build by default, as some C3 library dependencies (CBLAS) can induce small differences in our regression suite.
This capability is also being used as a prototype to explore model-based versus method-based specification of stochastic expansions. While the model specification is stand-alone, it currently requires a corresponding method specification to exercise the model, which can be a generic UQ strategy such as surrogate_based_uq method or a sampling method. The intent is to migrate function train, polynomial chaos, and stochastic collocation toward model-only specifications that can then be employed in any surrogate/emulator context.

Examples

```dakota
model,
  id_model = 'FT'
  surrogate global function_train
  start_order = 2
  start_rank = 2 kick_rank = 2  max_rank = 10
  adapt_rank
  dace_method_pointer = 'SAMPLING'
```

See Also

These keywords may also be of interest:

- function_train
- multilevel_function_train
- multifidelity_function_train

7.3.3.2.5.1 regression_type

- Keywords Area
- model
- surrogate
- global
- function_train
- regression_type

Type of solver for forming function train approximations by regression

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**
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<tr>
<th>Required/Optional Required</th>
<th>Description of Group Regression Type (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td></td>
<td></td>
<td>ls</td>
<td>Use least squares solver for forming function train approximations by regression</td>
</tr>
<tr>
<td></td>
<td></td>
<td>rls2</td>
<td>Use regularized regression solver for forming function train approximations</td>
</tr>
</tbody>
</table>

**Description**

Function train approximations are formed based on regression for the set of coefficients described at `function_train`. Solver options include least squares and regularized regression, where the latter penalizes high-order terms to mitigate overfitting.

**Default Behavior**

The default regression solver is least squares.

7.3.3.2.5.2 ls

- Keywords Area
- model
- surrogate
- global
- function_train
- regression_type
- ls

Use least squares solver for forming function train approximations by regression

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Least squares is the standard solver for forming function train approximations based on regression. This option does not include regularization.

7.3.3.2.5.3 rls2

- Keywords Area
- model
- surrogate
Use regularized regression solver for forming function train approximations

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional Required</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>l2_penalty</td>
<td>Penalty value applied in regularized regression solver for function train approximations</td>
</tr>
</tbody>
</table>

Description

RLS2 is a regularized least squares solver that applies an $L2$ penalty to mitigate overfitting. It includes the option to specify a penalty parameter.

See Also

These keywords may also be of interest:

- l2_penalty

7.3.3.2.5.4 l2_penalty

- Keywords Area
- model
- surrogate
- global
- function_train
- regression_type
- rls2
- l2_penalty

Penalty value applied in regularized regression solver for function train approximations
Specification

**Alias:** none  
**Argument(s):** REAL

**Description**

Regularized regression applies a penalty to aspects of the recovered solution (e.g., cardinality, higher-order oscillatory terms, et al.) to mitigate overfitting. This specification provides the scalar penalty parameter (its value is not currently adapted or cross-validated).

**Default Behavior**

If this penalty parameter is not specified, then we adopt the internal C3 default.

**See Also**

These keywords may also be of interest:

- rls2

7.3.3.2.5.5 `max_solver_iterations`

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 100

**Description**

When using an iterative polynomial coefficient estimation approach, e.g., cross-validation-based solvers, limits the maximum iterations in the coefficient solver.

7.3.3.2.5.6 `max_cross_iterations`
7.3 model

- function_train
- max_cross_iterations

Maximum number of iterations for cross-approximation during a rank adaptation.

Specification

Alias: none
Argument(s): INTEGER
Default: 1

Description

The number of iterations for the cross-approximation algorithm is limited within the adapt_rank procedure.

See Also

These keywords may also be of interest:

- adapt_rank

7.3.3.2.5.7 solver_tolerance

- Keywords Area
- model
- surrogate
- global
- function_train
- solver_tolerance

Convergence tolerance for the optimizer used during the regression solve.

Specification

Alias: none
Argument(s): REAL
Default: 1.0e-10

Description

Defines the function value and function gradient tolerance for optimizer used solving the regression problem for the functional tensor train coefficients.
7.3.3.2.5.8 response_scaling

• Keywords Area
• model
• surrogate
• global
• function_train
• response_scaling

Perform bounds-scaling on response values prior to surrogate emulation

Specification

Alias: none
Argument(s): none
Default: use original data

Description

Particularly in multifidelity methods based on regression solutions at each level of a model hierarchy, scaling can be an issue since the magnitude of discrepancy data may decay rapidly and regression solvers may employ absolute tolerances in places.

By activating response_scaling, the set of response data used for each recovery is scaled to [0,1] based on the minimum value and range of the data set. This prevents loss of accuracy due to imbalances in scale across model levels.

See Also

These keywords may also be of interest:

• multifidelity_polynomial_chaos
• multifidelity_function_train

7.3.3.2.5.9 tensor_grid

• Keywords Area
• model
• surrogate
• global
• function_train
• tensor_grid

Use sub-sampled tensor-product quadrature points to build a polynomial chaos expansion.
Specification

Alias: none
Argument(s): none
Default: regression with random sample set

Description

The collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

7.3.3.2.5.10 rounding_tolerance

- Keywords Area
- model
- surrogate
- global
- function_train
- rounding_tolerance

An accuracy tolerance that is used to guide rounding during rank adaptation.

Specification

Alias: none
Argument(s): REAL
Default: 1.e-10

Description

The rounding tolerance is used to measure how closely one can approximate a functional tensor train representation at a step of the optimization procedure using one of lower ranks. If this tolerance is small, then less compression is achieved but greater accuracy is obtained. If this tolerance is big, greater compression is achieved at the expense of accuracy.

Default Behavior

If this tolerance is not specified, then we adopt the internal C3 default.

See Also

These keywords may also be of interest:

- adapt_rank
- arithmetic_tolerance
7.3.3.2.5.11 arithmetic_tolerance

A secondary rounding tolerance used for post-processing

**Specification**

*Alias*: none  
*Argument(s)*: REAL  
*Default*: 1.e-10

**Description**

After forming one or more function train approximations, this tolerance is used for rolling up final results, in particular, combining multilevel/multifidelity expansions and forming products for higher-order moment estimation. It is generally intended as a looser tolerance than rounding_tolerance.  

**Default Behavior**

If this tolerance is not specified, then we adopt the internal C3 default.

**See Also**

These keywords may also be of interest:

- rounding_tolerance

7.3.3.2.5.12 start_order

(Initial) polynomial order of each univariate function within the functional tensor train.
Specification

Alias: order
Argument(s): INTEGER
Default: 2
Child Keywords:
### dimension_preference

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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>dimension_ preference</td>
<td>A set of weights specifying the relative importance of each uncertain variable (dimension)</td>
</tr>
</tbody>
</table>

#### Description

This specifies the initial polynomial order of each univariate function that makes up the functional tensor train decomposition. In the case of adapt_order, polynomial order starts from this value and is then advanced; its maximum value is limited by max_order and increments are controlled by kick_order.

#### Default Behavior

Default is 2.

#### See Also

These keywords may also be of interest:

- adapt_order
- kick_order
- max_order

#### Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** isotropic grids

#### Description

A set of weights specifying the relative importance of each uncertain variable (dimension). Using this specification leads to anisotropic integrations with differing refinement levels for different random dimensions.
See Also

These keywords may also be of interest:

- sobol
- decay

7.3.3.2.5.14 adapt_order

- Keywords Area
- model
- surrogate
- global
- function_train
- adapt_order

Activate adaptive procedure for determining the best basis order

Specification

Alias: none
Argument(s): none
Default: false

Description

This option activates a cross validation procedure to select the best basis order for representing the current set of data. It searches from start_order to max_order in increments of kick_order.

Default Behavior

No cross validation for basis order.

Examples

This example shows specification of an order adaptation starting at order 2, incrementing by 2, and limited at order 6.

```plaintext
model,
  id_model = 'FT'
surrogate global function_train
  start_rank = 4
  adapt_order start_order = 2 kick_order = 2 max_order = 6
  solver_tolerance = 1e-12
  rounding_tolerance = 1e-12
dace_method_pointer = 'SAMPLING'
```

Note that adapt_rank and adapt_order can either be combined or used separately.
See Also

These keywords may also be of interest:

- `start_order`
- `kick_order`
- `max_order`

### 7.3.3.2.5.15 kick_order

- Keywords Area
- model
- surrogate
- global
- function_train
- kick_order

Increment used when adapting the basis order in function train methods.

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 1

**Description**

Used in combination with `adapt_order, kick_order` defines the size of the increment that is used during the cross validation process, within the range of `start_order` and `max_order`.

**Default Behavior**

Default is 1.

**See Also**

These keywords may also be of interest:

- `adapt_order`

### 7.3.3.2.5.16 max_order

- Keywords Area
- model
- surrogate
- global
- function_train
- max_order

Maximum polynomial order of each univariate function within the functional tensor train.
Specification

Alias: none

Argument(s): INTEGER
Default: 4

Description

This specification is inactive for current adaptation options.
In future adaptation options (i.e., cross-approximation adaptation), this will specify the maximum order that can be obtained during adaptation, where polynomial order will start from $start\_order$ and be limited by $max\_order$.

Default Behavior

Max order is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- start_order

7.3.3.2.5.17 max_cv_order_candidates

- Keywords Area
- model
- surrogate
- global
- function_train
- max_cv_order_candidates

Limit the number of cross-validation candidates for basis order

Specification

Alias: none

Argument(s): INTEGER
Default: ushort max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.
See Also

These keywords may also be of interest:

- increment_max_order
- increment_max_rank_order
- polynomial_chaos
- multifidelity_polynomial_chaos
- function_train
- multifidelity_function_train

7.3.3.2.5.18 start_rank

The initial rank used for the starting point during a rank adaptation.

Specification

Alias: rank
Argument(s): INTEGER
Default: 2

Description

The adapt_rank procedure increments the rank on each iteration, starting from the rank defined by start_rank

Default Behavior
Default is 2.

See Also

These keywords may also be of interest:

- adapt_rank
7.3 model

7.3.3.2.5.19 adapt_rank

- Keywords Area
- model
- surrogate
- global
- function_train
- adapt_rank

Activate adaptive procedure for determining best rank representation

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** false

**Description**

The adaptive algorithm proceeds as follows:

1. Start from rank `start_rank` and form an approximation

2. Adapt the current approximation by searching for a solution with lower rank that achieves L2 accuracy within epsilon tolerance of the reference.

3. If a lower rank solution is found with comparable accuracy, then stop. If not, increase the rank by an amount specified by `kick_rank`.

4. Return to step 2 and continue until either `max_rank` is reached or a converged rank (rank less than current reference with comparable accuracy) is found.

**Default Behavior**

No cross validation for rank.

**Examples**

This example shows specification of a rank adaptation starting at rank 2, incrementing by 2, and limited at rank 10.

```plaintext
model,
  id_model = 'FT'
  surrogate global function_train
  start_order = 5
  adapt_rank start_rank = 2 kick_rank = 2 max_rank = 10
  solver_tolerance = 1e-12
  rounding_tolerance = 1e-12
  dace_method_pointer = 'SAMPLING'
```

Note that `adapt_rank` and `adapt_order` can either be combined or used separately.
See Also

These keywords may also be of interest:

- start_rank
- kick_rank
- max_rank

7.3.3.2.5.20 kick_rank

- Keywords Area
- model
- surrogate
- global
- function_train
- kick_rank

The increment in rank employed during each iteration of the rank adaptation.

Specification

Alias: none

Argument(s): INTEGER

Default: 2

Description

The adapt_rank procedure increments the rank on each iteration, using an increment defined by kick_rank.

Default Behavior

Default is 1.

See Also

These keywords may also be of interest:

- adapt_rank

7.3.3.2.5.21 max_rank

- Keywords Area
- model
- surrogate
- global
- function_train
- max_rank

Limits the maximum rank that is explored during a rank adaptation.
7.3 model

Specification

Alias: none
Argument(s): INTEGER
Default: 3

Description

The adapt_rank procedure increments the rank on each iteration, and the maximum value is limited by max_rank

Default Behavior
Max rank is unbounded (adaptive sweep will stop when CV error increases).

See Also

These keywords may also be of interest:

- adapt_rank

7.3.3.2.5.22 max_cv_rank_candidates

- Keywords Area
- model
- surrogate
- global
- function_train
- max_cv_rank_candidates

Limit the number of cross-validation candidates for rank

Specification

Alias: none
Argument(s): INTEGER
Default: size_t max

Description

When generating refinement candidates by advancing the upper bound used for a cross-validation range, the number of cross-validation can eventually become excessive and hamper algorithm progress. This control limits the maximum number of candidates, which when active, will increase the lower bound for this range as the upper bound is advanced.

See Also

These keywords may also be of interest:
Artificial neural network model

Specification

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

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<tr>
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<td>max_nodes</td>
<td></td>
<td>Maximum number of hidden layer nodes</td>
</tr>
<tr>
<td>Optional</td>
<td>range</td>
<td></td>
<td>Range for neural network random weights</td>
</tr>
<tr>
<td>Optional</td>
<td>random_weight</td>
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<td>(Inactive) Random weight control</td>
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<td>Optional</td>
<td>export_model</td>
<td></td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
<tr>
<td>Optional</td>
<td>import_model</td>
<td></td>
<td>Import surrogate model from archive file</td>
</tr>
</tbody>
</table>

**Description**

Dakota’s artificial neural network surrogate is a stochastic layered perceptron network, with a single hidden layer. Weights for the input layer are chosen randomly, while those in the hidden layer are estimated from data using a variant of the Zimmerman direct training approach[96].
This typically yields lower training cost than traditional neural networks, yet good out-of-sample performance. This is helpful in surrogate-based optimization and optimization under uncertainty, where multiple surrogates may be repeatedly constructed during the optimization process, e.g., a surrogate per response function, and a new surrogate for each optimization iteration.

The neural network is a non-parametric surface fitting method. Thus, along with Kriging (Gaussian Process) and MARS, it can be used to model data trends that have slope discontinuities as well as multiple maxima and minima. However, unlike Kriging, the neural network surrogate is not guaranteed to interpolate the data from which it was constructed.

This surrogate can be constructed from fewer than \( n_{\text{quad}} \) data points, however, it is a good rule of thumb to use at least \( n_{\text{quad}} \) data points when possible.

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

**Theory**

The form of the neural network model is

\[
\hat{f}(x) \approx \tanh \left \{ A_1 \tanh \left ( A_0^T x + \theta_0^T \right ) + \theta_1 \right \}
\]

where \( x \) is the evaluation point in \( n \)-dimensional parameter space; the terms \( A_0, \theta_0 \) are the random input layer weight matrix and bias vector, respectively; and \( A_1, \theta_1 \) are a weight vector and bias scalar, respectively, estimated from training data. These coefficients are analogous to the polynomial coefficients obtained from regression to training data. The neural network uses a cross validation-based orthogonal matching pursuit solver to determine the optimal number of nodes and to solve for the weights and offsets.

7.3.3.2.6.1 max_nodes

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **neural_network**
- **max_nodes**

Maximum number of hidden layer nodes

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** nodes

**Argument(s):** INTEGER

**Default:** numTrainingData - 1
Description

Limits the maximum number of hidden layer nodes in the neural network model. The default is to use one less node than the number of available training data points yielding a fully-determined linear least squares problem. However, reducing the number of nodes can help reduce overfitting and more importantly, can drastically reduce surrogate construction time when building from a large data set. (Historically, Dakota limited the number of nodes to 100.)

The keyword `max_nodes` provides an upper bound. Dakota’s orthogonal matching pursuit algorithm may further reduce the effective number of nodes in the final model to achieve better generalization to unseen points.

7.3.3.2.6.2 range

- Keywords Area
- model
- surrogate
- global
- neural_network
- range

Range for neural network random weights

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): REAL

Description

Controls the range of the input layer random weights in the neural network model. The default range is 2.0, resulting in weights in (-1, 1). These weights are applied after the training inputs have been scaled into [-0.8, 0.8].

7.3.3.2.6.3 random_weight

- Keywords Area
- model
- surrogate
- global
- neural_network
- random_weight

(Inactive) Random weight control
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): INTEGER

Description

This option is not currently in use and is likely to be removed

7.3.3.2.6.4 export_model

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
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<td>filename_prefix</td>
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<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required</td>
<td>formats</td>
<td></td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>
Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- text_archive - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- binary_archive - Binary, machine-readable archive for re-import or use with the surfpack executable
- algebraic_file - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- algebraic_console - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword gaussian_process surfpack)
- Artificial neural network (keyword neural_network)
- Radial basis Functions (keyword radial_basis)
- Polynomial (keyword polynomial)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword mars) and moving least squares (keyword moving_least_squares) models, only text_archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

Default Behavior

No export.

Expected Output

Output depends on selected format; see the formats specification.

Additional Discussion

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.6.5 filename_prefix

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive
```

7.3.3.2.6.6 formats

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- formats

Formats for surrogate model export

Topics

This keyword is related to the topics:

- surrogate_models
### Specification

**Alias:** none  
**Argument(s):** none  

### Child Keywords:

<table>
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</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>text_archive</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_file</td>
<td>algebraic_file</td>
<td>Export surrogate model in algebraic format to a file</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_console</td>
<td>algebraic_console</td>
<td>Export surrogate model in algebraic format to the console</td>
</tr>
</tbody>
</table>

### Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted.  
See `export_model` and the entries for the format selection keywords for further information.

#### 7.3.3.2.6.7 `text_archive`

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format

### Topics

This keyword is related to the topics:

- surrogate_models
Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named \{prefix\}.{response_descriptor}.txt. Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.sps, in which ‘sps’ stands for Surfpack surrogate. See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.6.8 binary_archive

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named \{prefix\}.{response_descriptor}.bin.
Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.bsps, in which 'bsps' stands for binary Surfpack surrogate. See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.6.9 algebraic_file

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- formats
- algebraic_file

Export surrogate model in algebraic format to a file

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to a file in a human-readable "algebraic" format. The file is named using the pattern {prefix}.{response_descriptor}.alg. See filename_prefix for further information about exported surrogate file naming. The file contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the file matches exactly the output written to the console when algebraic_console is specified.
7.3.3.6.10 algebraic_console

- Keywords Area
- model
- surrogate
- global
- neural_network
- export_model
- formats
- algebraic_console

Export surrogate model in algebraic format to the console

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to the console (screen, or output file if Dakota was run using the -o option) in a human-readable "algebraic" format. The output contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the screen for the exported model matches exactly the output written to file when algebraic_file is specified. Use of algebraic_file is preferred over algebraic_console, which exists largely to provide a measure of backward compatibility.

7.3.3.6.11 import_model

- Keywords Area
- model
- surrogate
- global
- neural_network
- import_model

Import surrogate model from archive file
Specification

Alias: none
Argument(s): none
Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
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<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
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<td>Surrogate Import Format (Group 1)</td>
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<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
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</table>

Description

Imports a surrogate model from a file in binary_archive or text_archive format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords filename_prefix and binary_archive or text_archive.

Default Behavior

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through dace_method_pointer.

Usage Tips

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing export_model and its child keywords to those needed for import_model. Any other keywords such as specifying a dace_iterator or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed initial_state values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the surfpack binary or the experimental dakota.surrogates Python module, it is untested and not the primary use case for this capability.
Examples

The following model block exports a Gaussian process after it's built:

```dakota
model
  id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
  metrics 'root_mean_squared'
cross_validation folds 5
export_model
  filename_prefix = 'gp_export'
  formats = binary_archive
  export_approx_points_file 'gp_values.dat'
  export_approx_variance_file 'gp_variance.dat'

And this model block imports it using a nearly identical model specification:

```dakota
model
  id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
experimental_gaussian_process
  metrics 'root_mean_squared'
cross_validation folds 5
import_model
  filename_prefix = 'gp_export'
binary_archive
  export_approx_points_file 'gp_values.dat'
  export_approx_variance_file 'gp_variance.dat'

7.3.3.2.6.12 filename_prefix

- Keywords Area
- model
- surrogate
- global
- neural_network
- import_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern \( \text{prefix}.\{\text{response_descriptor}\}.\text{extension} \). This filename_prefix keyword is used to supply the prefix portion of the pattern.

The response_descriptor portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns \( \text{my_surrogate}.\{\text{response_descriptor}\}.\text{alg} \) (for the algebraic files) and \( \text{my_surrogate}.\{\text{response_descriptor}\}.\text{bsps} \) (for the binary files).

```
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
formats
  algebraic_file
  binary_archive
```

7.3.2.6.13 text_archive

- Keywords Area
- model
- surrogate
- global
- neural_network
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias**: none

**Argument(s)**: none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named \( \text{prefix}.\{\text{response_descriptor}\}.\text{txt} \).
Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.2.6.14 binar Archive

• Keywords Area
• model
• surrogate
• global
• neural_network
• import_model
• binary_archive

 Surrogate model binary archive file format

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named `{prefix}.{response_descriptor}.bin`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Radial basis function (RBF) model

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

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<th>Description</th>
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<td>bases</td>
<td></td>
<td>Initial number of radial basis functions</td>
</tr>
<tr>
<td>Optional</td>
<td>max_pts</td>
<td></td>
<td>Maximum number of RBF CVT points</td>
</tr>
<tr>
<td>Optional</td>
<td>min_partition</td>
<td></td>
<td>(Inactive) Minimum RBF partition</td>
</tr>
<tr>
<td>Optional</td>
<td>max_subsets</td>
<td></td>
<td>Number of trial RBF subsets</td>
</tr>
<tr>
<td>Optional</td>
<td>export_model</td>
<td></td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
<tr>
<td>Optional</td>
<td>import_model</td>
<td></td>
<td>Import surrogate model from archive file</td>
</tr>
</tbody>
</table>

**Description**

Radial basis functions \( \phi \) are functions whose value typically depends on the distance from a center point, called the centroid, \( c \).

The surrogate model approximation comprises a sum of \( K \) weighted radial basis functions:

\[
\hat{f}(x) = \sum_{k=1}^{K} w_k \phi(\|x - c_k\|)
\]

These basis functions take many forms, but Gaussian kernels or splines are most common. The Dakota implementation uses a Gaussian radial basis function. The weights are determined via a linear least squares solution approach. See[70] for more details.
Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

7.3.3.2.7.1 bases

- Keywords Area
- model
- surrogate
- global
- radial_basis
- bases

Initial number of radial basis functions

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): INTEGER

Description

Initial number of radial basis functions. The default value is the smaller of the number of training points and 100.

7.3.3.2.7.2 max_pts

- Keywords Area
- model
- surrogate
- global
- radial_basis
- max_pts

Maximum number of RBF CVT points

Topics

This keyword is related to the topics:

- surrogate_models
Specification

**Alias:** none  
**Argument(s):** INTEGER

Description

Maximum number of CVT points to use in generating each RBF center. Default is computing centroid of each. Defaults to $10 \times \text{(bases)}$. Reducing this will reduce model build time.

### 7.3.3.2.7.3 min_partition

- Keywords Area
- model
- surrogate
- global
- radial_basis
- min_partition

(Inactive) Minimum RBF partition

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none  
**Argument(s):** INTEGER

Description

This option currently has no effect and will likely be removed.

### 7.3.3.2.7.4 max_subsets

- Keywords Area
- model
- surrogate
- global
- radial_basis
- max_subsets

Number of trial RBF subsets
7.3 model

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): INTEGER

Description

Number of passes to take to identify the best subset of basis functions to use. Defaults to the smaller of $3 \times \text{(bases)}$ and 100.

7.3.3.2.7.5 export_model

- Keywords Area
  - model
  - surrogate
  - global
  - radial_basis
  - export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

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<td>Required</td>
<td>filename_prefix</td>
<td>formats</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td></td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>
Description

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive** - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive** - Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file** - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console** - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only **text_archive** and **binary_archive** formats may be used.

Currently, no other surrogate models can be exported.

**Default Behavior**

No export.

**Expected Output**

Output depends on selected format; see the formats specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Topics
This keyword is related to the topics:

- surrogate_models

Specification

Alias: none
Argument(s): STRING
Default: exported_surrogate

Description
Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples
This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
    filename_prefix = 'my_surrogate'
    formats
        algebraic_file
        binary_archive
```

7.3.3.2.7.7 formats

- Keywords Area
- model
- surrogate
- global
- radial_basis
- export_model
- formats

Formats for surrogate model export

Topics
This keyword is related to the topics:

- surrogate_models
**Specification**

**Alias:** none  
**Argument(s):** none  

**Child Keywords:**

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<th>Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td>text_archive</td>
<td></td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td></td>
<td>Surrogate model binary archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_file</td>
<td></td>
<td>Export surrogate model in algebraic format to a file</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_console</td>
<td></td>
<td>Export surrogate model in algebraic format to the console</td>
</tr>
</tbody>
</table>

**Description**

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See `export_model` and the entries for the format selection keywords for further information.

7.3.3.2.7.8 text_archive  

- Keywords Area
- model
- surrogate
- global
- radial_basis
- export_model
- formats
- text_archive

Surrogate model plain-text archive file format

**Topics**

This keyword is related to the topics:

- surrogate_models
7.3 model

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.
Experimental surrogates will export to (import from) text archives named {prefix}.{response_descriptor}.txt.
Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.sps, in which 'sps' stands for Surfpack surrogate.
See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.7.9 binary_archive

• Keywords Area
• model
• surrogate
• global
• radial_basis
• export_model
• formats
• binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.
Experimental surrogates will export to (import from) binary archives named {prefix}.{response_descriptor}.bin.
Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.7.10 algebraic_file

- Keywords Area
- model
- surrogate
- global
- radial_basis
- export_model
- formats
- algebraic_file

Export surrogate model in algebraic format to a file

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to a file in a human-readable "algebraic" format. The file is named using the pattern `{prefix}.{response_descriptor}.alg`. See `filename_prefix` for further information about exported surrogate file naming. The file contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the file matches exactly the output written to the console when `algebraic_console` is specified.
7.3.3.2.7.11 algebraic_console

• Keywords Area
• model
• surrogate
• global
• radial_basis
• export_model
• formats
• algebraic_console

Export surrogate model in algebraic format to the console

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to the console (screen, or output file if Dakota was run using the -o option) in a human-readable “algebraic” format. The output contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the screen for the exported model matches exactly the output written to file when algebraic_file is specified. Use of algebraic_file is preferred over algebraic_console, which exists largely to provide a measure of backward compatibility.

7.3.3.2.7.12 import_model

• Keywords Area
• model
• surrogate
• global
• radial_basis
• import_model

Import surrogate model from archive file
Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Surrogate Import Format (Group 1)</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Imports a surrogate model from a file in binary_archive or text_archive format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords filename_prefix and binary_archive or text_archive.

Default Behavior

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through dace_method_pointer.

Usage Tips

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing export_model and its child keywords to those needed for import_model. Any other keywords such as specifying a dace_iterator or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed initial_state values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the surfpack binary or the experimental dakota.surrogates Python module, it is untested and not the primary use case for this capability.
Examples

The following model block exports a Gaussian process after it’s built:

```plaintext
model
  id_model = 'SURR_M'
  surrogate global
    dace_method_pointer = 'DACE'
    experimental_gaussian_process
      metrics 'root_mean_squared'
      cross_validation folds 5
      export_model
        filename_prefix = 'gp_export'
        formats = binary_archive
        export_approx_points_file 'gp_values.dat'
        export_approx_variance_file 'gp_variance.dat'

And this model block imports it using a nearly identical model specification:

```plaintext
model
  id_model = 'SURR_M'
  surrogate global
    dace_method_pointer = 'DACE'
    experimental_gaussian_process
      metrics 'root_mean_squared'
      cross_validation folds 5
      import_model
        filename_prefix = 'gp_export'
        binary_archive
        export_approx_points_file 'gp_values.dat'
        export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.7.13 filename_prefix

- Keywords Area
- model
- surrogate
- global
- radial_basis
- import_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models

Specification

- Alias: none
- Argument(s): STRING
- Default: exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
forms
  algebraic_file
  binary_archive
```

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias**: none

**Argument(s)**: none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. 
Other (Surfpack) surrogates will export/import using files named \( \text{prefix}.\text{response\_descriptor}.\text{sps} \), in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

**Examples**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.7.15  binary_archive

- **Keywords Area**
  - model
  - surrogate
  - global
  - radial_basis
  - import_model
  - binary_archive

Surrogate model binary archive file format

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named \( \text{prefix}.\text{response\_descriptor}.\text{bin} \).

Other (Surfpack) surrogates will export/import using files named \( \text{prefix}.\text{response\_descriptor}.\text{bsps} \), in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

**Examples**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
7.3.3.2.8 polynomial

• Keywords Area
• model
• surrogate
• global
• polynomial

Polynomial surrogate model

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Use a linear polynomial or trend function</td>
<td></td>
</tr>
<tr>
<td></td>
<td>quadratic</td>
<td>Use a quadratic polynomial or trend function</td>
<td></td>
</tr>
<tr>
<td></td>
<td>cubic</td>
<td>Use a cubic polynomial</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>export_model</td>
<td>Exports surrogate model in user-specified format(s)</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>import_model</td>
<td>Import surrogate model from archive file</td>
<td></td>
</tr>
</tbody>
</table>

Description

Linear, quadratic, and cubic polynomial surrogate models are available in Dakota. The utility of the simple polynomial models stems from two sources:

• over a small portion of the parameter space, a low-order polynomial model is often an accurate approximation to the true data trends
• the least-squares procedure provides a surface fit that smooths out noise in the data.

Local surrogate-based optimization methods (surrogate_based_local) are often successful when using polynomial models, particularly quadratic models. However, a polynomial surface fit may not be the best choice for modeling data trends globally over the entire parameter space, unless it is known a priori that the true data trends are close to linear, quadratic, or cubic. See[65] for more information on polynomial models.
Known Issue: When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

Theory

The form of the linear polynomial model is

\[ \hat{f}(x) \approx c_0 + \sum_{i=1}^{n} c_i x_i \]

the form of the quadratic polynomial model is:

\[ \hat{f}(x) \approx c_0 + \sum_{i=1}^{n} c_i x_i + \sum_{i=1}^{n} \sum_{j \geq i} c_{ij} x_i x_j \]

and the form of the cubic polynomial model is:

\[ \hat{f}(x) \approx c_0 + \sum_{i=1}^{n} c_i x_i + \sum_{i=1}^{n} \sum_{j \geq i} c_{ij} x_i x_j + \sum_{i=1}^{n} \sum_{j \geq i} \sum_{k \geq j} c_{ijk} x_i x_j x_k \]

In all of the polynomial models, \( \hat{f}(x) \) is the response of the polynomial model; the \( x_i, x_j, x_k \) terms are the components of the \( n \)-dimensional design parameter values; the \( c_0, c_i, c_{ij}, c_{ijk} \) terms are the polynomial coefficients, and \( n \) is the number of design parameters. The number of coefficients, \( n_c \), depends on the order of polynomial model and the number of design parameters. For the linear polynomial:

\[ n_{c,linear} = n + 1 \]

for the quadratic polynomial:

\[ n_{c,quad} = \frac{(n + 1)(n + 2)}{2} \]

and for the cubic polynomial:

\[ n_{c,cubic} = \frac{n^3 + 6n^2 + 11n + 6}{6} \]

There must be at least \( n_c \) data samples in order to form a fully determined linear system and solve for the polynomial coefficients. In Dakota, a least-squares approach involving a singular value decomposition numerical method is applied to solve the linear system.

7.3.3.2.8.1 basis_order

- Keywords Area
- model
- surrogate
- global
- polynomial
- basis_order
Specification

**Alias:** none

**Argument(s):** INTEGER

Description

The polynomial order for the polynomial regression model (default = 2).

### 7.3.3.2.8.2 linear

- Keywords Area
- model
- surrogate
- global
- polynomial
- linear

Use a linear polynomial or trend function

Specification

**Alias:** none

**Argument(s):** none

Description

See parent page

### 7.3.3.2.8.3 quadratic

- Keywords Area
- model
- surrogate
- global
- polynomial
- quadratic

Use a quadratic polynomial or trend function

Specification

**Alias:** none

**Argument(s):** none
7.3 model

Description

See parent page

7.3.3.2.8.4 cubic

- Keywords Area
- model
- surrogate
- global
- polynomial
- cubic

Use a cubic polynomial

Specification

Alias: none

Argument(s): none

Description

See parent page

7.3.3.2.8.5 export_model

- Keywords Area
- model
- surrogate
- global
- polynomial
- export_model

Exports surrogate model in user-specified format(s)

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:
Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the formats specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive**: Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive**: Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file**: Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console**: Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword `gaussian_process surfpack`)
- Artificial neural network (keyword `neural_network`)
- Radial basis Functions (keyword `radial_basis`)
- Polynomial (keyword `polynomial`)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword `mars`) and moving least squares (keyword `moving_least_squares`) models, only text_-archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

**Default Behavior**

No export.

**Expected Output**

Output depends on selected format; see the `formats` specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.8.6  filename_prefix

- Keywords Area
- model
- surrogate
7.3 model

- global
- polynomial
- export_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern {prefix}.{response_descriptor}.{extension}. This filename_prefix keyword is used to supply the prefix portion of the pattern.

The response_descriptor portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns my_surrogate.{response_descriptor}.alg (for the algebraic files) and my_surrogate.{response_descriptor}.bsps (for the binary files).

```
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
  formats
    algebraic_file
    binary_archive

7.3.3.2.8.7 formats
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• formats

Formats for surrogate model export

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td>text_archive</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>binary_archive</td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_file</td>
<td>algebraic_file</td>
<td>Export surrogate model in algebraic format to a file</td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_console</td>
<td>algebraic_console</td>
<td>Export surrogate model in algebraic format to the console</td>
</tr>
</tbody>
</table>

Description

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See export_model and the entries for the format selection keywords for further information.

7.3.3.2.8.8 text_archive

• Keywords Area
• model
• surrogate
• global
• polynomial
• export_model
• formats
• text_archive

Surrogate model plain-text archive file format
Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named \{prefix\}.{response_descriptor}.txt. Other (Surfpack) surrogates will export/import using files named \{prefix\}.{response_descriptor}.sps, in which 'sps' stands for Surfpack surrogate. See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.8.9  binary_archive

- Keywords Area
- model
- surrogate
- global
- polynomial
- export_model
- formats
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none
Description

When specified, the surrogate model will be exported (or imported) using a binary archive format. Experimental surrogates will export to (import from) binary archives named \( \text{prefix}.\{\text{response}\_\text{descriptor}\}.\text{bin} \). Other (Surfpack) surrogates will export/import using files named \( \text{prefix}.\{\text{response}\_\text{descriptor}\}.\text{bsps} \), in which ‘bsps’ stands for binary Surfpack surrogate.

See \text{filename\_prefix} for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.8.10 algebraic\_file

- Keywords Area
- model
- surrogate
- global
- polynomial
- export\_model
- formats
- algebraic\_file

Export surrogate model in algebraic format to a file

Topics

This keyword is related to the topics:

- surrogate\_models

Specification

\text{Alias}: \text{none}

\text{Argument(s)}: \text{none}

Description

After the surrogate model has been built, Dakota will export it to a file in a human-readable “algebraic” format. The file is named using the pattern \( \text{prefix}.\{\text{response}\_\text{descriptor}\}.\text{alg} \). See \text{filename\_prefix} for further information about exported surrogate file naming. The file contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the file matches exactly the output written to the console when \text{algebraic\_console} is specified.
7.3.3.2.8.11 algebraic_console

• Keywords Area
• model
• surrogate
• global
• polynomial
• export_model
• formats
• algebraic_console

Export surrogate model in algebraic format to the console

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): none

Description

After the surrogate model has been built, Dakota will export it to the console (screen, or output file if Dakota was run using the -o option) in a human-readable “algebraic” format. The output contains sufficient information for the user to (re)construct and evaluate the model outside of Dakota.

Expected Output

The format depends on the type of surrogate model, but in general will include a LaTeX-like representation of the analytic form of the model to aid tool development, all needed model hyperparameters, and headers describing the shape or dimension of the provided data.

The output written to the screen for the exported model matches exactly the output written to file when algebraic_file is specified. Use of algebraic_file is preferred over algebraic_console, which exists largely to provide a measure of backward compatibility.

7.3.3.2.8.12 import_model

• Keywords Area
• model
• surrogate
• global
• polynomial
• import_model

Import surrogate model from archive file
Specification

Alias: none

Argument(s): none

Child Keywords:

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<thead>
<tr>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>filename_prefix</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Surrogate Import Format (Group 1)</td>
<td>text_archive</td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>binary_archive</td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

Description

Imports a surrogate model from a file in binary_archive or text_archive format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords filename_prefix and binary_archive or text_archive.

Default Behavior

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through dace_method_pointer.

Usage Tips

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing export_model and its child keywords to those needed for import_model. Any other keywords such as specifying a dace_iterator or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed initial_state values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the surfpack binary or the experimental dakota.surrogates Python module, it is untested and not the primary use case for this capability.
Examples

The following model block exports a Gaussian process after it’s built:

```plaintext
model
  id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
eperimental_gaussian_process
  metrics 'root_mean_squared'
  cross_validation folds 5
export_model
  filename_prefix = 'gp_export'
  formats = binary_archive
  export_approx_points_file 'gp_values.dat'
  export_approx_variance_file 'gp_variance.dat'
```

And this model block imports it using a nearly identical model specification:

```plaintext
model
  id_model = 'SURR_M'
surrogate global
dace_method_pointer = 'DACE'
eperimental_gaussian_process
  metrics 'root_mean_squared'
  cross_validation folds 5
import_model
  filename_prefix = 'gp_export'
  binary_archive
  export_approx_points_file 'gp_values.dat'
  export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.8.13 filename_prefix

- Keywords Area
- model
- surrogate
- global
- polynomial
- import_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate
Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.

Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```dakota
surrogate global gaussian_process surfpack
export_model
  filename_prefix = 'my_surrogate'
formats
  algebraic_file
  binary_archive
```

7.3.3.2.8.14 text_archive

- Keywords Area
- model
- surrogate
- global
- polynomial
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias**: none

**Argument(s)**: none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.

Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. 
Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which 'sps' stands for Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.8.15 binary_archive

- Keywords Area
- model
- surrogate
- global
- polynomial
- import_model
- binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named `{prefix}.{response_descriptor}.bin`.

Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.bsps`, in which 'bsps' stands for binary Surfpack surrogate.

See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
Use a deterministic polynomial surrogate

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

<table>
<thead>
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<tbody>
<tr>
<td>Required</td>
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<td>basis_order</td>
<td>Total degree of the polynomial basis</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>options_file</td>
<td>Filename for a YAML file that specifies polynomial surrogate options</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>export_model</td>
<td>Exports surrogate model in user-specified format(s)</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>import_model</td>
<td>Import surrogate model from archive file</td>
</tr>
</tbody>
</table>

**Description**

Compute a polynomial surrogate for a specified total order by least-squares regression. This model is implemented in Dakota's surrogates module and is considered experimental.

7.3.3.2.9.1 basis_order

- Keywords Area
- model
- surrogate
- global
- experimental_polynomial
- basis_order

Total degree of the polynomial basis
7.3 model

Specification

Alias: none
Argument(s): INTEGER

Description

This keyword's argument specifies the terms in the polynomial according to a total order scheme.

7.3.3.2.9.2 options_file

- Keywords Area
- model
- surrogate
- global
- experimental_polynomial
- options_file

Filename for a YAML file that specifies polynomial surrogate options

Specification

Alias: none
Argument(s): STRING
Default: no advanced options file

Description

The YAML file contains configuration options that are used to populate a Teuchos ParameterList used by the polynomial that will override other keyword-specified parameters. Missing options in the YAML file are set to default values.

Examples

# FILE: polynomial_options.yaml

Polynomial Parameters:
- max degree: 3
- scaler type: none
- reduced basis: false
- p-norm: 1.0
- regression solver type: SVD
- standardize response: false
- verbosity: 1

7.3.3.2.9.3 export_model

- Keywords Area
- model
- surrogate
Exports surrogate model in user-specified format(s)

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tr>
<td>Required</td>
<td>filename_prefix</td>
<td>formats</td>
<td>User-customizable portion of exported/imported surrogate model filenames</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Formats for surrogate model export</td>
</tr>
</tbody>
</table>

**Description**

Export the surrogate for later re-import to Dakota or evaluation using the surfpack executable (bin/surfpack) or a user-developed tool. Export format is controlled using the **formats** specification. Four formats are available in Dakota; however, not all have been enabled for all surrogates.

The four formats are:

- **text_archive** - Plain-text, machine-readable archive for re-import or use with the surfpack executable
- **binary_archive** - Binary, machine-readable archive for re-import or use with the surfpack executable
- **algebraic_file** - Plain-text, human-readable file intended for use with user-created tools; not compatible with Dakota or the surfpack executable
- **algebraic_console** - Print the model in algebraic format to the screen; not compatible with Dakota or the surfpack executable

These global surrogates can be exported in all four formats:

- Gaussian process (keyword **gaussian_process** surfpack)
- Artificial neural network (keyword **neural_network**)
- Radial basis Functions (keyword **radial_basis**)
• Polynomial (keyword polynomial)

However, for experimental Gaussian Process and polynomial models as well as Multivariate Adaptive Regression Spline (keyword mars) and moving least squares (keyword moving_least_squares) models, only text-archive and binary_archive formats may be used.

Currently, no other surrogate models can be exported.

**Default Behavior**

No export.

**Expected Output**

Output depends on selected format; see the formats specification.

**Additional Discussion**

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.9.4 filename_prefix

- Keywords Area
- model
- surrogate
- global
- experimental_polynomial
- export_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

**Topics**

This keyword is related to the topics:

- surrogate_models

**Specification**

*Alias:* none

*Argument(s):* STRING

*Default:* exported_surrogate

**Description**

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern {prefix}.{response_descriptor}.{extension}. This filename_prefix keyword is used to supply the prefix portion of the pattern.

The response_descriptor portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.
Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```plaintext
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive
```

### 7.3.3.2.9.5 formats

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **experimental_polynomial**
- **export_model**
- **formats**

Formats for surrogate model export

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** none  
**Argument(s):** none

**Child Keywords:**

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<th>Description</th>
<th>Dakota Keyword</th>
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</tr>
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<tr>
<td></td>
<td><strong>Optional</strong></td>
<td>text_archive</td>
<td></td>
<td>Surrogate model plain-text archive file format</td>
</tr>
<tr>
<td></td>
<td><strong>Optional</strong></td>
<td>binary_archive</td>
<td></td>
<td>Surrogate model binary archive file format</td>
</tr>
</tbody>
</table>

**Description**

Select from among the 2-4 available export formats available for this surrogate. Multiple selections are permitted. See `export_model` and the entries for the format selection keywords for further information.
Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** none  
**Argument(s):** none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format.  
Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`.  
Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.  
See `filename_prefix` for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
• formats
• binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none
Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.
Experimental surrogates will export to (import from) binary archives named \( \text{prefix} \cdot \text{response_descriptor} \cdot \text{bin} \).
Other (Surfpack) surrogates will export/import using files named \( \text{prefix} \cdot \text{response_descriptor} \cdot \text{bsps} \), in which ‘bsps’ stands for binary Surfpack surrogate.
See \text{filename_prefix} for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.9.8 import_model

• Keywords Area
• model
• surrogate
• global
• experimental_polynomial
• import_model

Import surrogate model from archive file

Specification

Alias: none
Argument(s): none
Child Keywords:
**Description**

Imports a surrogate model from a file in `binary_archive` or `text_archive` format, typically instead of constructing it from generated or imported data. Importing can result in significant time savings with some models, such as Gaussian processes. The file from which to import is further specified with the child keywords `filename_prefix` and `binary_archive` or `text_archive`.

**Default Behavior**

When used in the context of an iteratively adapted or rebuilt surrogate, the imported model will only replace the initial surrogate model. Subsequent surrogate model builds require imported build points or a design of experiments method specified through `dace_method_pointer`.

**Usage Tips**

When importing a surrogate model, it is crucial that the global surrogate model part of the Dakota input file be identical for export and import, except for changing `export_model` and its child keywords to those needed for `import_model`. Any other keywords such as specifying a `dace_iterator` or imported points must remain intact to satisfy internal surrogate constructor requirements.

All variables over which the surrogate model was built (typically the active variables) when exported must be present in the Dakota study that imports it. The variables in the importing study will be matched by descriptor to those in the surrogate model. This allows the order and type of variables to change between export and import, for example to fix unimportant parameters as state variables even if they were previously active design variables.

For example, suppose the original Dakota study for the surrogate build had active uniform uncertain variables: a, b, c, d, and inactive state variables s, t, and the surrogate gets built over a, b, c, d. The follow-on study could perform design optimization over active design variables b and d, with inactive state variables a, c, s, t, held at fixed `initial_state` values. In this case the values of a, b, c, d will get mapped to the surrogate and s, t dropped when evaluating it. So it is important the state values for s and t haven’t changed in some way that invalidates the surrogate.

While it might be possible to import surrogates externally-generated using the `surfpack` binary or the experimental `dakota.surrogates` Python module, it is untested and not the primary use case for this capability.

**Examples**

The following model block exports a Gaussian process after it’s built:

```dakota
model
    id_model = 'SURR_M'
    surrogate global
        dace_method_pointer = 'DACE'
        experimental gaussian process
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
And this model block imports it using a nearly identical model specification:

```dakota
model
  id_model = 'SURR_M'
  surrogate global
    dace_method_pointer = 'DACE'
    experimental_gaussian_process
      metrics 'root_mean_squared'
      cross_validation folds 5
  import_model
    filename_prefix = 'gp_export'
    binary_archive
    export_approx_points_file 'gp_values.dat'
    export_approx_variance_file 'gp_variance.dat'
```

7.3.3.2.9.9 filename_prefix

- Keywords Area
- model
- surrogate
- global
- experimental_polynomial
- import_model
- filename_prefix

User-customizable portion of exported/imported surrogate model filenames

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): STRING

Default: exported_surrogate

Description

Dakota surrogate model files are exported and imported using one file per response, per requested format. The files are named using the pattern `{prefix}.{response_descriptor}.{extension}`. This `filename_prefix` keyword is used to supply the prefix portion of the pattern.

The `response_descriptor` portion of the pattern is filled in using the response descriptors provided by the user (or, if none are specified, descriptors automatically generated by Dakota). Extension is a three or four letter string that depends on the format.
Examples

This input snippet directs Dakota to write one algebraic format file and one binary archive file for each response. The names of the files will follow the patterns `my_surrogate.{response_descriptor}.alg` (for the algebraic files) and `my_surrogate.{response_descriptor}.bsps` (for the binary files).

```
surrogate global gaussian_process surfpack
export_model
   filename_prefix = 'my_surrogate'
   formats
      algebraic_file
      binary_archive
```

7.3.3.2.9.10 text_archive

- Keywords Area
- model
- surrogate
- global
- experimental_polynomial
- import_model
- text_archive

Surrogate model plain-text archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a plain-text archive format. Experimental surrogates will export to (import from) text archives named `{prefix}.{response_descriptor}.txt`. Other (Surfpack) surrogates will export/import using files named `{prefix}.{response_descriptor}.sps`, in which ‘sps’ stands for Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.
7.3.3.2.9.11 binary_archive

- Keywords Area
  - model
  - surrogate
  - global
  - experimental_polynomial
  - import_model
  - binary_archive

Surrogate model binary archive file format

Topics

This keyword is related to the topics:

- surrogate_models

Specification

Alias: none

Argument(s): none

Description

When specified, the surrogate model will be exported (or imported) using a binary archive format.

Experimental surrogates will export to (import from) binary archives named {prefix}.{response_descriptor}.bin.

Other (Surfpack) surrogates will export/import using files named {prefix}.{response_descriptor}.bsps, in which 'bsps' stands for binary Surfpack surrogate.

See filename_prefix for further information about surrogate file naming.

Examples

The Dakota examples include a demonstration of using the surfpack executable with an exported model file.

7.3.3.2.10 domain_decomposition

- Keywords Area
  - model
  - surrogate
  - global
  - domain_decomposition

Piecewise Domain Decomposition for Global Surrogate Models
## Specification

**Alias:** none  
**Argument(s):** none

### Child Keywords:

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<td>Optional</td>
<td></td>
<td>discontinuity_detection</td>
<td>Optional Discontinuity Detection Capability for the Piecewise Decomposition Option of Global Surrogates</td>
</tr>
</tbody>
</table>

### Description

Typical regression techniques use all available sample points to build continuous approximations the underlying function.

An alternative option is to use piecewise decomposition to locally approximate the function at some point using a few sample points from its neighborhood only. This option currently supports Polynomial Regression, Gaussian Process (GP) Interpolation, and Radial Basis Functions (RBF) Regression. It requires a decomposition cell type (currently set to be Voronoi cells). Optional parameters are: the number of layers of neighbors used to solve the regression problem (default is one layer), and an optional discontinuity detection capability (identified by a user-input jump or gradient threshold).

The method can also make use of the gradient and Hessian information, if available. The user needs to specify the keyword `user_derivatives`.

### 7.3.3.10.1 cell_type

- Keywords Area
- model
- surrogate
- global
- domain_decomposition
• **cell_type**

Type of the Geometric Cells Used for the Piecewise Decomposition Option of Global Surrogates

**Specification**

**Alias:** none  
**Argument(s):** STRING

**Description**

The piecewise decomposition option for global surrogates is used to locally approximate a function at some point using a few sample points from its neighborhood. This option requires a decomposition cell type that can vary from structured grid boxes, to polygonal Voronoi cells. Currently, this option only supports Voronoi cells.

### 7.3.3.2.10.2 support_layers

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **domain_decomposition**
- **support_layers**

Optional Number of Support Layers for the Piecewise Decomposition Option of Global Surrogates

**Specification**

**Alias:** none  
**Argument(s):** INTEGER

**Description**

The piecewise decomposition option for global surrogates is used to locally approximate a function at some point using a few sample points from its neighborhood. The neighborhood of a cell is parameterized via a number of (support layers). The default value is set to one layer of neighbors (cells that share direct edges with the cell under study). One more support layer would include the neighbors of that cell's neighbors, and so on.

### 7.3.3.2.10.3 discontinuity_detection

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **domain_decomposition**
Optional Discontinuity Detection Capability for the Piecewise Decomposition Option of Global Surrogates

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<td><strong>jump_threshold</strong></td>
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<tr>
<td></td>
<td></td>
<td><strong>gradient_threshold</strong></td>
<td>Gradient Threshold Parameter of the Optional Discontinuity Detection Capability for the Piecewise Decomposition Option of Global Surrogates</td>
</tr>
</tbody>
</table>

**Description**

The piecewise decomposition option for global surrogates is used to locally approximate a function at some point using a few sample points from its neighborhood.

The domain decomposition algorithm supports an optional discontinuity detection capability where seeds across a user-input discontinuity threshold are not considered neighbors when building the approximate connectivity Delaunay graph. Alternatively, the domain is split into patches that trap discontinuities between them. This capability is specified by either jump or gradient threshold values in the input spec.

7.3.3.2.10.4  jump_threshold

- **Keywords Area**
  - model
  - surrogate
  - global
  - domain_decomposition
  - discontinuity_detection
• jump_threshold

Gradient Threshold Parameter of the Optional Discontinuity Detection Capability for the Piecewise Decomposition Option of Global Surrogates

Specification

Alias: none
Argument(s): REAL

Description

The piecewise decomposition option for global surrogates is used to locally approximate a function at some point using a few sample points from its neighborhood.

The domain decomposition algorithm supports an optional discontinuity detection capability where seeds across a user-input discontinuity threshold are not considered neighbors when building the approximate connectivity Delaunay graph. Alternatively, the domain is split into patches that trap discontinuities between them. This capability can be specified using a jump threshold value in the input spec.
7.3.3.2.11 total_points

- Keywords Area
- model
- surrogate
- global
- total_points

Specified number of training points

Specification

**Alias**: none

**Argument(s)**: INTEGER

**Default**: recommended_points

Description

See parent page.

7.3.3.2.12 minimum_points

- Keywords Area
- model
- surrogate
- global
- minimum_points

Construct surrogate with minimum number of points

Specification

**Alias**: none

**Argument(s)**: none

Description

The minimum is d+1, for d input dimensions, except for polynomials. See parent page.

7.3.3.2.13 recommended_points

- Keywords Area
- model
- surrogate
- global
- recommended_points

Construct surrogate with recommended number of points
Specification

**Alias:** none

**Argument(s):** none

Description

This is the default option. It requires $5d$ build points for $d$ input dimensions, except for polynomial models. See parent page.

7.3.3.2.14 dace_method_pointer

- **Keywords Area**
- model
- surrogate
- global
- dace_method_pointer

Specify a method to gather training data

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** no design of experiments data

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td></td>
<td>auto_refinement</td>
<td>Experimental auto-refinement of surrogate model</td>
</tr>
</tbody>
</table>

Description

The number of training points and the sources are specified on `global`, as well as the number of new training points required.

New training points are gathered by running the “truth” model using the method specified by `dace_method_pointer`. The DACE method will only be invoked if it has new samples to perform, and if new samples are required and no DACE iterator has been provided, an error will result.

The `dace_method_pointer` points to design of experiments method block used to generate truth model data.

Permissible methods include: Monte Carlo (random) sampling, Latin hypercube sampling, orthogonal array sampling, central composite design sampling, and Box-Behnken sampling.
Note that the number of samples specified in the method block may be overwritten, if the requested number of samples is less than `minimum_points`.

### 7.3.3.2.14.1 auto_refinement

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **dace_method_pointer**
- **auto_refinement**

Experimental auto-refinement of surrogate model

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no refinement

**Child Keywords:**

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<th>Description of Group</th>
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<tr>
<td>Optional</td>
<td>max_iterations</td>
<td></td>
<td>Number of iterations allowed for optimizers and adaptive UQ methods</td>
</tr>
<tr>
<td>Optional</td>
<td>max_function_-_evaluations</td>
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<td>Number of function evaluations allowed for optimizers</td>
</tr>
<tr>
<td>Optional</td>
<td>convergence_-_tolerance</td>
<td></td>
<td>Cross-validation threshold for surrogate convergence</td>
</tr>
</tbody>
</table>
### Description

(Experimental option) Automatically refine the surrogate model until desired cross-validation quality is achieved. Refinement is accomplished by iteratively adding more data to the training set until the cross-validation convergence_tolerance is achieved, or max_function_evaluations or max_iterations is exceeded.

The amount of new training data that is incorporated each iteration is specified in the DACE method that is referred to by the model's dace_method_pointer. See refinement_samples for more information.

#### 7.3.3.2.14.2 max_iterations

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **dace_method_pointer**
- **auto_refinement**
- **max_iterations**

Number of iterations allowed for optimizers and adaptive UQ methods

#### Topics

This keyword is related to the topics:

- **method_independent_controls**

#### Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 100

#### Description

The maximum number of iterations is used as a stopping criterion for optimizers and some adaptive UQ methods. If it has not reached any other stopping criteria first, the method will stop after it has performed max_iterations iterations. In cases where there are multiple nested iterations to be controlled, max_iterations generally controls the outer-most context.

**Default Behavior**

Default value is 100.
See Also

These keywords may also be of interest:

- max_function_evaluations

7.3.3.2.14.3 max_function_evaluations

- Keywords Area
- model
- surrogate
- global
- dace_method_pointer
- auto_refinement
- max_function_evaluations

Number of function evaluations allowed for optimizers

Topics

This keyword is related to the topics:

- method_independent_controls

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** 1000

**Description**

The maximum number of function evaluations is used as a stopping criterion for optimizers. If it has not reached any other stopping criteria first, the optimizer will stop after it has performed max_function_evaluations evaluations. See also max_iterations.

Some optimizers (e.g. ncsu_direct) may run past this limit in the course of an iteration step that began before max_function_evaluations was exceeded.

**Default Behavior**

Default value is 1000.

7.3.3.2.14.4 convergence_tolerance

- Keywords Area
- model
- surrogate
- global
- dace_method_pointer
Cross-validation threshold for surrogate convergence

**Specification**

**Alias:** none  
**Argument(s):** REAL  
**Default:** 1.0e-4

**Description**

The surrogate model will be refined until the selected `cross_validation_metric` falls below this convergence tolerance.

**Default Behavior**  The default is 1e-4.

7.3.3.2.14.5  `soft_convergence_limit`

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **dace_method_pointer**
- **auto_refinement**
- **soft_convergence_limit**

Maximum number of iterations without improvement in cross-validation

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 0

**Description**

The purpose of this control is to cease auto-refinement when improvement in the cross-validation score appears to have stalled or occurs too slowly.

A rolling average of improvement in the cross-validation score is computed over `soft_convergence_limit` iterations. Auto-refinement is halted if the average becomes smaller than the `convergence_tolerance`. The average is computed but ignored until the number of iterations has exceeded `soft_convergence_limit`. The default setting of 0 disables soft convergence.
7.3.3.2.14.6  cross_validation_metric

- Keywords Area
- model
- surrogate
- global
- dace_method_pointer
- auto_refinement
- cross_validation_metric

Choice of error metric to satisfy

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** root-mean-squared error

**Child Keywords:**

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<th>Required/Optional</th>
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<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>folds</td>
<td>Number of cross validation folds</td>
</tr>
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</table>

**Description**

The cross-validation score that the auto-refinement routine seeks to satisfy is based on an error metric. Any of the surrogate diagnostic metrics may be used. These include:

- root_mean_squared
- mean_abs
- rsquared
- sum_squared
- mean_squared
- sum_abs
- max_abs

**Default Behavior**  The default metric is the root mean squared error ("root_mean_squared").

7.3.3.2.14.7  folds

- Keywords Area
- model
- surrogate
- global
• dace_method_pointer
• auto_refinement
• cross_validation_metric
• folds

Number of cross validation folds

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** 10

**Description**

Number of folds (partitions) of the training data to use in cross validation (default 10).

---

### 7.3.3.2.15 actual_model_pointer

- Keywords Area
- model
- surrogate
- global
- actual_model_pointer

A surrogate model pointer that guides a method to whether it should use a surrogate model or compute truth function evaluations

**Specification**

**Alias:** truth_model_pointer

**Argument(s):** STRING

**Description**

Dakota methods use global surrogate models to compute surrogate function approximations. They also need to know the true function evaluations. A global surrogate model now must have an actual_model_pointer keyword to decide for the method whether to evaluate the global surrogate model, or compute the true function evaluations if actual_model_pointer = ‘TRUTH’.

---

### 7.3.3.2.16 reuse_points

- Keywords Area
- model
- surrogate
- global
- reuse_points

Surrogate model training data reuse control
Topics

This keyword is related to the topics:

- surrogate_models

Specification

**Alias:** reuse_samples

**Argument(s):** none

**Default:** all for import; none otherwise

**Child Keywords:**

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<td></td>
<td>all</td>
<td>region Option for reuse_points</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>none Option for reuse_points</td>
</tr>
</tbody>
</table>

**Description**

Dakota’s global surrogate methods rely on training data, which can either come from evaluation of a “truth” model, which is generated by the method specified with dace_method_pointer, from a file of existing training data, identified by import_build_points_file, or both.

The `reuse_points` keyword controls the amount of training data used in building a surrogate model, either initially, or during iterative rebuild, as in surrogate-based optimization. If import_build_points_file is specified, `reuse_points` controls how the file contents are used. If used during iterative rebuild, it controls what data from previous surrogate builds is reused in building the current model.

- **all** (default for file import) - use all points in the file or available from previous builds
- **region** - use only the points falling in the current trust region (see surrogate_based_local)
- **none** (default when no import) - ignore the contents of the file or previous build points, and gather new training data using the specified DACE method

7.3.3.2.16.1 all

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **reuse_points**
- **all**

Option for reuse_points
Specification

**Alias:** none

**Argument(s):** none

Description

This is described on the parent page.

7.3.3.2.16.2 region

- Keywords Area
- model
- surrogate
- global
- reuse_points
- region

Option for reuse_points

Specification

**Alias:** none

**Argument(s):** none

Description

This is described on the parent page.

7.3.3.2.16.3 none

- Keywords Area
- model
- surrogate
- global
- reuse_points
- none

Option for reuse_points

Specification

**Alias:** none

**Argument(s):** none
Description

This is described on the parent page.

7.3.3.2.17 import_build_points_file

- Keywords Area
- model
- surrogate
- global
- import_build_points_file

File containing points you wish to use to build a surrogate

Specification

**Alias:** import_points_file samples_file

**Argument(s):** STRING

**Default:** no point import from a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional (Choose One)</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>custom_annotated</td>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td>freeform</td>
<td>Selects freeform file format</td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td></td>
<td>Import only active variables from tabular data file</td>
</tr>
</tbody>
</table>

**Description**

The `import_build_points_file` allows the user to specify a file that contains a list of points and truth model responses used to construct a surrogate model. These can be used by all methods that (explicitly, e.g. surrogate-based optimization, or implicitly, e.g. efficient global optimization) operate on a surrogate. In particular, these points and responses are used in place of truth model evaluations to construct the initial surrogate. When used to construct surrogate models or emulators these are often called build points or training data.

**Default Behavior**

By default, methods do not import points from a file.

**Usage Tips**

Dakota parses input files without regard to whitespace, but the `import_build_points_file` must be in one of three formats:

- annotated (default)
• custom_annotated
• freeform

Examples

method
  polynomial_chaos
  expansion_order = 4
  import_build_points_file = 'dakota_uq_rosenbrock_pce_import.annot.pts.dat'

7.3.3.2.17.1 custom_annotated

• Keywords Area
• model
• surrogate
• global
• import_build_points_file
• custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>header</td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>eval_id</td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>interface_id</td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```
environment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```
%eval_id x1  x2   obj_fn nln_ineq_con_1 nln_ineq_con_2
1     0.9 1.1  0.0002 0.26 0.76
2 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.3.3.2.17.2 header

- Keywords Area
  - model
  - surrogate
  - global
  - import_build_points_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Child Keywords:
Table:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>use_variable_labels</td>
<td>Validate/use variable labels from tabular file header</td>
</tr>
</tbody>
</table>

Description

See description of parent `custom_annotated`

7.3.2.17.3  use_variable_labels

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- custom_annotated
- header
- use_variable_labels

Validate/use variable labels from tabular file header

Specification

Alias: none

Argument(s): none

Description

When importing global surrogate training data (or challenge evaluation points) from a tabular data file containing a header (annotated or custom_annotated header), this keyword toggles validation of variable labels present in the header row.

The labels are validated against the descriptors of the variables being imported to. If the tabular file labels can be rearranged to match the expected labels, the columns in the data file will be reordered on read to match the Dakota variable order. If the read labels are not a permutation of, nor equal to, the expected labels, an error will result.

Default Behavior

When not specified, variable labels will be read, but not strictly enforced or reordered. A warning will be issued if the variable labels are not as expected, and guidance offered if they can be permuted to match expectations.

Expected Output

Console output will be generated for any warnings, as well as to indicate whether any variable reordering is taking place.

Usage Tips

The use of this keyword is recommended when importing header-annotated tabular data files where the variables are appropriately labeled. Tabular files do not always contain response labels, so no attempt is made to disambiguate variable from response labels. The variable labels must appear contiguously in the header after any leading column IDs such as 'eval_id' or 'interface'.
Examples

This example enforces variable labels for both build and challenge points

```python
model
    id_model = 'SURR'
surrogate global
    polynomial quadratic
    import_build_points_file = 'dakota_surrogate_import.unc_fixedothers.dat'
    annotated use_variable_labels
challenge_points_file = 'dakota_surrogate_import.dat'
    annotated use_variable_labels
```

7.3.3.2.17.4 eval_id

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent custom_annotated

7.3.3.2.17.5 interface_id

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none
See description of parent custom_annotated

7.3.3.2.17.6 annotated

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>use_variable_labels</td>
<td>Validate/use variable labels from tabular file header</td>
</tr>
</tbody>
</table>

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

%eval_id interface x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1 NO_ID 0.9 1.1 0.0002 0.26 0.76
2 NO_ID 0.90009 1.1 0.0001996404857 0.2601620081 0.759955
3 NO_ID 0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...```

7.3.3.2.17.7 use_variable_labels

• Keywords Area

• model

• surrogate

• global

• import_build_points_file

• annotated

• use_variable_labels

Validate/use variable labels from tabular file header

Specification

Alias: none

Argument(s): none

Description

When importing global surrogate training data (or challenge evaluation points) from a tabular data file containing a header (annotated or custom_annotated header), this keyword toggles validation of variable labels present in the header row.

The labels are validated against the descriptors of the variables being imported to. If the tabular file labels can be rearranged to match the expected labels, the columns in the data file will be reordered on read to match the Dakota variable order. If the read labels are not a permutation of, nor equal to, the expected labels, an error will result.

Default Behavior

When not specified, variable labels will be read, but not strictly enforced or reordered. A warning will be issued if the variable labels are not as expected, and guidance offered if they can be permuted to match expectations.
Expected Output

Console output will be generated for any warnings, as well as to indicate whether any variable reordering is taking place.

Usage Tips

The use of this keyword is recommended when importing header-annotated tabular data files where the variables are appropriately labeled. Tabular files do not always contain response labels, so no attempt is made to disambiguate variable from response labels. The variable labels must appear contiguously in the header after any leading column IDs such as 'eval_id' or 'interface'.

Examples

This example enforces variable labels for both build and challenge points:

```model
id_model = 'SURR'
surrogate global
  polynomial quadratic
import_build_points_file = 'dakota_surrogate_import.unc_fixedothers.dat'
  annotated use_variable_labels
challenge_points_file = 'dakota_surrogate_import.dat'
  annotated use_variable_labels
```

7.3.3.2.17.8 freeform

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Default: annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.
Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.3.3.2.17.9 active_only

- Keywords Area
- model
- surrogate
- global
- import_build_points_file
- active_only

Import only active variables from tabular data file

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.3.3.2.18 `export_approx_points_file`

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **export_approx_points_file**

Output file for surrogate model value evaluations

Specification

**Alias:** `export_points_file`

**Argument(s):** STRING

**Default:** no point export to a file

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Optional/Choose</strong></td>
<td>Tabular Format (Group 1)</td>
<td><code>custom.annotated</code></td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td><strong>One</strong></td>
<td></td>
<td><code>annotated</code></td>
<td>Selects annotated tabular file format</td>
</tr>
<tr>
<td></td>
<td></td>
<td><code>freeform</code></td>
<td>Selects freeform file format</td>
</tr>
</tbody>
</table>

Description

Specifies a file in which the points (input variable values) at which the surrogate model is evaluated and corresponding response values computed by the surrogate model will be written. The response values are the surrogate’s predicted approximation to the truth model responses at those points.

Usage Tips

Dakota exports tabular data in one of three formats:

- `annotated` *(default)*
- `custom.annotated`
- `freeform`
7.3.3.2.18.1 custom_annotated

- **Keywords Area**
  - model
  - surrogate
  - global
  - export_approx_points_file
  - custom_annotated

Selects custom-annotated tabular file format

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

**Description**

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether header row, eval_id column, and interface_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

**Usage Tips**
Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
environment
  tabular_data
  tabular_data_file = 'dakota_summary.dat'
  custom_annotated header eval_id

Resulting tabular file:

```
%eval_id x1 x2 obj_fn nln_ineq_con_1 nln_ineq_con_2
1  0.9  1.1  0.0002  0.26  0.76
2  0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3  0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

Enable header row in custom-annotated tabular file

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

See description of parent custom_annotated
7.3.3.2.18.3  eval_id

- Keywords Area
- model
- surrogate
- global
- export_approx_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated

7.3.3.2.18.4  interface_id

- Keywords Area
- model
- surrogate
- global
- export_approx_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Description

See description of parent custom_annotated
7.3.3.2.18.5  annotated

- Keywords Area
- model
- surrogate
- global
- export_approx_points_file
- annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

Default Behavior

By default, Dakota imports and exports tabular files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- To specify pre-Dakota 6.1 tabular format, which did not include interface_id, specify custom_annotated header eval_id

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.
Examples

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
  tabular_data_file = 'dakota_summary.dat'
annotated
```

Resulting tabular file:

<table>
<thead>
<tr>
<th>%eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nl_n_ineq_con_1</th>
<th>nl_n_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.9009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

7.3.3.2.18.6 freeform

• Keywords Area
  • model
  • surrogate
  • global
  • export_approx_points_file

Selects freeform file format

Topics

This keyword is related to the topics:

• file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.

**Default Behavior**

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

**Usage Tips**
• Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

• When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

• In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

• Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    freeform
```

Resulting tabular file:

```
0.9 1.1 0.0002 0.26 0.76
0.90009 1.1 0.0001996404857 0.2601620081 0.759955
0.89991 1.1 0.0002003604863 0.2598380081 0.760045
...
```

7.3.3.2.19 use_derivatives

• Keywords Area

• model

• surrogate

• global

• use_derivatives

Use derivative data to construct surrogate models

Specification

Alias: none

Argument(s): none

Default: use function values only

Description

The use_derivatives flag specifies that any available derivative information should be used in global approximation builds, for those global surrogate types that support it (currently, polynomial regression and the Surfpack Gaussian process).

However, it’s use with Surfpack Gaussian process is not recommended.
Correction approaches for surrogate models

Specification

Alias: none

Argument(s): none

Default: no surrogate correction

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Correction Order (Group 1)</td>
<td>zeroth_order</td>
<td>Specify that truth values must be matched.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>first_order</td>
<td>Specify that truth values and gradients must be matched.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>second_order</td>
<td>Specify that truth values, gradients and Hessians must be matched.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Required (Choose One)</th>
<th>Correction Type (Group 2)</th>
<th>additive</th>
<th>Additive correction factor for local surrogate accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>multiplicative</td>
<td>Multiplicative correction factor for local surrogate accuracy.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>combined</td>
<td>Multipoint correction for a hierarchical surrogate</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

Some of the surrogate model types support the use of correction factors that improve the local accuracy of the surrogate models.

The correction specification specifies that the approximation will be corrected to match truth data, either matching truth values in the case of zeroth_order matching, matching truth values and gradients in the case of first_order matching, or matching truth values, gradients, and Hessians in the case of second_order matching. For additive and multiplicative corrections, the correction is local in that the truth data is matched at a single point, typically the center of the approximation region. The additive correction adds a scalar offset (zeroth_order), a linear function (first_order), or a quadratic function (second_order) to the approximation to match the truth data at the point, and the multiplicative correction multiplies the approximation by a scalar (zeroth_order), a linear function (first_order), or a quadratic function (second_order) to match the truth data at the point. The additive first_order case is due to [59] and the multiplicative first_order case is commonly known as beta correction [41]. For the combined correction, the use of both additive and multiplicative corrections allows the satisfaction of an additional matching condition, typically the truth function values at the previous correction point (e.g., the center of the previous trust region). The combined correction is then a multipoint correction, as opposed to the local additive and multiplicative corrections. Each of these correction capabilities is described in detail in [24].

The correction factors force the surrogate models to match the true function values and possibly true function derivatives at the center point of each trust region. Currently, Dakota supports either zeroth-, first-, or second-order accurate correction methods, each of which can be applied using either an additive, multiplicative, or combined correction function. For each of these correction approaches, the correction is applied to the surrogate model and the corrected model is then interfaced with whatever algorithm is being employed. The default behavior is that no correction factor is applied.

The simplest correction approaches are those that enforce consistency in function values between the surrogate and original models at a single point in parameter space through use of a simple scalar offset or scaling applied to the surrogate model. First-order corrections such as the first-order multiplicative correction (also known as beta correction [14]) and the first-order additive correction [59] also enforce consistency in the gradients and provide a much more substantial correction capability that is sufficient for ensuring provable convergence in SBO algorithms. SBO convergence rates can be further accelerated through the use of second-order corrections which also enforce consistency in the Hessians [24], where the second-order information may involve analytic, finite-difference, or quasi-Newton Hessians.

Correcting surrogate models with additive corrections involves

\[ f_{hi}(x) = f_{lo}(x) + \alpha(x) \]  

where multifidelity notation has been adopted for clarity. For multiplicative approaches, corrections take the form

\[ f_{hi}(x) = f_{lo}(x) \beta(x) \]  

where, for local corrections, \( \alpha(x) \) and \( \beta(x) \) are first or second-order Taylor series approximations to the exact correction functions:

\[ \alpha(x) = A(x_c) + \nabla A(x_c)^T (x - x_c) + \frac{1}{2} (x - x_c)^T \nabla^2 A(x_c) (x - x_c) \]  

\[ \beta(x) = B(x_c) + \nabla B(x_c)^T (x - x_c) + \frac{1}{2} (x - x_c)^T \nabla^2 B(x_c) (x - x_c) \]  

where the exact correction functions are

\[ A(x) = f_{hi}(x) - f_{lo}(x) \]  

\[ B(x) = \frac{f_{hi}(x)}{f_{lo}(x)} \]
Refer to[24] for additional details on the derivations. A combination of additive and multiplicative corrections can provide for additional flexibility in minimizing the impact of the correction away from the trust region center. In other words, both additive and multiplicative corrections can satisfy local consistency, but through the combination, global accuracy can be addressed as well. This involves a convex combination of the additive and multiplicative corrections:

\[ \hat{f}_{hi}(x) = \gamma \hat{f}_{hi}(\alpha(x)) + (1 - \gamma) \hat{f}_{hi}(\beta(x)) \]

where \( \gamma \) is calculated to satisfy an additional matching condition, such as matching values at the previous design iterate.

It should be noted that in both first order correction methods, the function \( \hat{f}(x) \) matches the function value and gradients of \( f_t(x) \) at \( x = x_c \). This property is necessary in proving that the first order-corrected SBO algorithms are provably convergent to a local minimum of \( f_t(x) \). However, the first order correction methods are significantly more expensive than the zeroth order correction methods, since the first order methods require computing both \( \nabla f_t(x_c) \) and \( \nabla f_s(x_c) \). When the SBO strategy is used with either of the zeroth order correction methods, or with no correction method, convergence is not guaranteed to a local minimum of \( f_t(x) \). That is, the SBO strategy becomes a heuristic optimization algorithm. From a mathematical point of view this is undesirable, but as a practical matter, the heuristic variants of SBO are often effective in finding local minima.

### Usage guidelines

- Both the additive zeroth_order and multiplicative zeroth_order correction methods are “free” since they use values of \( f_t(x_c) \) that are normally computed by the SBO strategy.

- The use of either the additive first_order method or the multiplicative first_order method does not necessarily improve the rate of convergence of the SBO algorithm.

- When using the first order correction methods, the gradient-related response keywords must be modified to allow either analytic or numerical gradients to be computed. This provides the gradient data needed to compute the correction function.

- For many computationally expensive engineering optimization problems, gradients often are too expensive to obtain or are discontinuous (or may not exist at all). In such cases the heuristic SBO algorithm has been an effective approach at identifying optimal designs[35].

### zeroth_order

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **correction**
- **zeroth_order**

Specify that truth values must be matched.

### Specification

**Alias**: none

**Argument(s)**: none
Description

The correction specification specifies that the approximation will be corrected to match truth data. The keyword `zeroth_order` matching ensures that truth values are matched.

7.3.3.2.20.2 first_order

- Keywords Area
- model
- surrogate
- global
- correction
- first_order

Specify that truth values and gradients must be matched.

Specification

Alias: none

Argument(s): none

Description

This correction specification specifies that the approximation will be corrected to match truth data. The keyword `first_order` matching ensures that truth values and gradients are matched.

7.3.3.2.20.3 second_order

- Keywords Area
- model
- surrogate
- global
- correction
- second_order

Specify that truth values, gradients and Hessians must be matched.

Specification

Alias: none

Argument(s): none

Description

The correction specification specifies that the approximation will be corrected to match truth data. The keyword `second_order` matching ensures that truth values, gradients and Hessians are matched.
7.3.3.20.4 additive

- Keywords Area
- model
- surrogate
- global
- correction
- additive

Additive correction factor for local surrogate accuracy

Specification

Alias: none
Argument(s): none

Description

Use an additive correction factor to improve the local accuracy of a surrogate.

7.3.3.20.5 multiplicative

- Keywords Area
- model
- surrogate
- global
- correction
- multiplicative

Multiplicative correction factor for local surrogate accuracy.

Specification

Alias: none
Argument(s): none

Description

Use a multiplicative correction factor to improve the local accuracy of a surrogate.
Multipoint correction for a hierarchical surrogate

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

For the combined correction, the use of both additive and multiplicative corrections allows the satisfaction of an additional matching condition, typically the truth function values at the previous correction point (e.g., the center of the previous trust region). The combined correction is then a multipoint correction, as opposed to the local additive and multiplicative corrections.

**7.3.3.2.21 metrics**

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **metrics**

Compute surrogate quality metrics

**Topics**

This keyword is related to the topics:

- **surrogate_models**

**Specification**

**Alias:** diagnostics

**Argument(s):** STRINGLIST

**Default:** No diagnostics

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>cross_validation</td>
<td>press</td>
<td>Leave-one-out cross validation</td>
</tr>
<tr>
<td>Optional</td>
<td>press</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Description

Diagnostic metrics assess the goodness of fit of a global surrogate to its training data. The default diagnostics are:

- root_mean_squared
- mean_abs
- rsquared

Additional available diagnostics include

- sum_squared
- mean_squared
- sum_abs
- max_abs

The keywords press and cross_validation further specify leave-one-out or k-fold cross validation, respectively, for all of the active metrics from above.

Usage Tips When specified, the metrics keyword must be followed by a list of quoted strings, each of which activates a metric.

Examples

This example input fragment constructs a quadratic polynomial surrogate and computes four metrics on the fit, both with and without 5-fold cross validation. (Also see dakota/share/dakota/test/dakota_surrogate_import.in for additional examples.)

```
model surrogate global polynomial quadratic
metrics = "root_mean_squared" "sum_abs" "mean_abs" "max_abs"
cross_validation folds = 5
```

Theory

Most of these diagnostics refer to some operation on the residuals (the difference between the surrogate model and the truth model at the data points upon which the surrogate is built).

For example, sum_squared refers to the sum of the squared residuals, and mean_abs refers to the mean of the absolute value of the residuals. rsquared refers to the R-squared value typically used in regression analysis (the proportion of the variability in the response that can be accounted for by the surrogate model). Care should be taken when interpreting metrics, for example, errors may be near zero for interpolatory models or rsquared may not be applicable for non-polynomial models.
• cross_validation

Perform k-fold cross validation

Topics

This keyword is related to the topics:

• surrogate_models

Specification

Alias: none

Argument(s): none

Default: No cross validation

Child Keywords:

<table>
<thead>
<tr>
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<th>Description of Group Partition Control (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional/Choose One</td>
<td></td>
<td>percent</td>
<td>Percent data per cross validation fold</td>
</tr>
</tbody>
</table>

Description

General k-fold cross validation may be performed by specifying cross_validation. The cross-validation statistics will be calculated for all metrics.

Cross validation may further specify:

• folds, the number of folds into which to divide the build data (between 2 and number of data points) or
• percent, the fraction of data (between 0 and 0.5) to use in each fold.

These will be adjusted as needed based on the number of available training points. The default number of folds k = 10, or 0.1
7.3 model

Specification

Alias: none
Argument(s): INTEGER
Default: 10

Description

Number of folds (partitions) of the training data to use in cross validation (default 10).

7.3.3.21.3 percent

• Keywords Area
• model
• surrogate
• global
• metrics
• cross_validation
• percent

Percent data per cross validation fold

Specification

Alias: none
Argument(s): REAL
Default: 0.1

Description

Percent of the training data to use in each cross validation fold (default 0.1).

7.3.3.21.4 press

• Keywords Area
• model
• surrogate
• global
• metrics
• press

Leave-one-out cross validation
Specification

Alias: none
Argument(s): none
Default: No PRESS cross validation

Description

Leave-one-out (PRESS) cross validation may be performed by specifying `press`. The cross-validation statistics will be calculated for all metrics.

7.3.3.22 import_challenge_points_file

- Keywords Area
- model
- surrogate
- global
- import_challenge_points_file

Datafile of points to assess surrogate quality

Specification

Alias: challenge_points_file
Argument(s): STRING
Default: no user challenge data

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format</td>
</tr>
<tr>
<td>annotated</td>
<td>Selects annotated tabular file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>freeform</td>
<td>Selects freeform file format</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>active_only</td>
<td>Import only active variables from tabular data file</td>
<td></td>
</tr>
</tbody>
</table>

Description

Specifies a data file containing variable and response (truth) values, in one of three formats:

- annotated (default)
- custom_annotated
The surrogate is evaluated at the points in the file, and the surrogate (approximate) responses are compared against the truth results from the file. All metrics specified with `metrics` will be computed for the challenge data.

7.3.3.2.22.1 custom_annotated

- Keywords Area
  - model
  - surrogate
  - global
  - import_challenge_points_file
  - custom_annotated

Selects custom-annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>eval_id</td>
<td></td>
<td>Enable evaluation ID column in custom-annotated tabular file</td>
</tr>
<tr>
<td>Optional</td>
<td>interface_id</td>
<td></td>
<td>Enable interface ID column in custom-annotated tabular file</td>
</tr>
</tbody>
</table>

Description

A custom-annotated tabular file is a whitespace-separated text file typically containing row data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well. Custom-annotated allows user options for whether `header row`, `eval_id column`, and `interface_id column` appear in the tabular file, thus bridging freeform and (fully) annotated.
Default Behavior

The annotated format is the default for tabular export/import. To control which header row and columns are in the input/output, specify custom_annotated, followed by options, in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a custom-annotated tabular file in Dakota 6.0 format, which contained only header and eval_id (no interface_id), and data for variables and responses. Input file fragment:

```plaintext
evironment
  tabular_data
    tabular_data_file = 'dakota_summary.dat'
    custom_annotated header eval_id
```

Resulting tabular file:

```plaintext
%eval_id  x1  x2  obj_fn nln_ineq_con_1 nln_ineq_con_2
1         0.9  1.1  0.0002  0.26   0.76
2         0.90009 1.1  0.0001996404857  0.2601620081  0.759955
3         0.89991 1.1  0.0002003604863  0.2598380081  0.760045
...
```

Enable header row in custom-annotated tabular file

Specification

Alias: none

Argument(s): none

Child Keywords:
**Description**

See description of parent custom.annotated

7.3.3.22.3 use_variable_labels

- Keywords Area
- model
- surrogate
- global
- import_challenge_points_file
- custom.annotated
- header
- use_variable_labels

Validate/use variable labels from tabular file header

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When importing global surrogate training data (or challenge evaluation points) from a tabular data file containing a header (annotated or custom.annotated header), this keyword toggles validation of variable labels present in the header row.

The labels are validated against the descriptors of the variables being imported to. If the tabular file labels can be rearranged to match the expected labels, the columns in the data file will be reordered on read to match the Dakota variable order. If the read labels are not a permutation of, nor equal to, the expected labels, an error will result.

**Default Behavior**

When not specified, variable labels will be read, but not strictly enforced or reordered. A warning will be issued if the variable labels are not as expected, and guidance offered if they can be permuted to match expectations.

**Expected Output**

Console output will be generated for any warnings, as well as to indicate whether any variable reordering is taking place.

**Usage Tips**

The use of this keyword is recommended when importing header-annotated tabular data files where the variables are appropriately labeled. Tabular files do not always contain response labels, so no attempt is made to disambiguate variable from response labels. The variable labels must appear contiguously in the header after any leading column IDs such as 'eval_id' or 'interface'.
Examples

This example enforces variable labels for both build and challenge points

```python
model
  id_model = 'SURR'
surrogate global
  polynomial quadratic
  import_build_points_file = 'dakota_surrogate_import.unc_fixedothers.dat'
  annotated use_variable_labels
challenge_points_file = 'dakota_surrogate_import.dat'
  annotated use_variable_labels
```

7.3.3.22.4 eval_id

- Keywords Area
- model
- surrogate
- global
- import_challenge_points_file
- custom_annotated
- eval_id

Enable evaluation ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none

Description

See description of parent custom_annotated

7.3.3.22.5 interface_id

- Keywords Area
- model
- surrogate
- global
- import_challenge_points_file
- custom_annotated
- interface_id

Enable interface ID column in custom-annotated tabular file

Specification

**Alias:** none

**Argument(s):** none
Description

See description of parent `custom_annotated`

7.3.3.22.6  annotated

- **Keywords Area**
  - model
  - surrogate
  - global
  - `import_challenge_points_file`
  - annotated

Selects annotated tabular file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td><code>use_variable_labels</code></td>
<td>Validate/use variable labels from tabular file header</td>
</tr>
</tbody>
</table>

**Description**

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. Each subsequent row contains an evaluation ID and interface ID, followed by data for variables, or variables followed by responses, depending on context.

**Default Behavior**

By default, Dakota imports and exports tabular files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**

- To specify pre-Dakota 6.1 tabular format, which did not include `interface_id`, specify `custom_annotated header eval_id`

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the *freeform* option.

**Examples**

Export an annotated top-level tabular data file containing a header row, leading eval_id and interface_id columns, and data for variables and responses. Input file fragment:

```plaintext
environment
tabular_data
tabular_data_file = 'dakota_summary.dat'
annotated

Resulting tabular file:

<table>
<thead>
<tr>
<th>eval_id</th>
<th>interface</th>
<th>x1</th>
<th>x2</th>
<th>obj_fn</th>
<th>nln_ineq_con_1</th>
<th>nln_ineq_con_2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NO_ID</td>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>2</td>
<td>NO_ID</td>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>3</td>
<td>NO_ID</td>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
<tr>
<td>...</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

7.3.3.2.22.7 *use_variable_labels*

- **Keywords Area**
- **model**
- **surrogate**
- **global**
- **import_challenge_points_file**
- **annotated**
- **use_variable_labels**

Validate/use variable labels from tabular file header

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

When importing global surrogate training data (or challenge evaluation points) from a tabular data file containing a header (*annotated* or *custom_annotated* header), this keyword toggles validation of variable labels present in the header row.

The labels are validated against the descriptors of the variables being imported to. If the tabular file labels can be rearranged to match the expected labels, the columns in the data file will be reordered on read to match the Dakota variable order. If the read labels are not a permutation of, nor equal to, the expected labels, an error will result.

**Default Behavior**

When not specified, variable labels will be read, but not strictly enforced or reordered. A warning will be issued if the variable labels are not as expected, and guidance offered if they can be permuted to match expectations.
7.3 model

Expected Output

Console output will be generated for any warnings, as well as to indicate whether any variable reordering is taking place.

Usage Tips

The use of this keyword is recommended when importing header-annotated tabular data files where the variables are appropriately labeled. Tabular files do not always contain response labels, so no attempt is made to disambiguate variable from response labels. The variable labels must appear contiguously in the header after any leading column IDs such as 'eval_id' or 'interface'.

Examples

This example enforces variable labels for both build and challenge points

```plaintext
model
  id_model = 'SURR'
  surrogate global
    polynomial quadratic
    import_build_points_file = 'dakota_surrogate_import.unc_fixedothers.dat'
    annotated use_variable_labels
    challenge_points_file = 'dakota_surrogate_import.dat'
    annotated use_variable_labels
```

7.3.3.22.8 freeform

- Keywords Area
- model
- surrogate
- global
- import_challenge_points_file
- freeform

Selects freeform file format

Topics

This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. Most commonly, each row contains data for variables, or variables followed by responses, though the format is used for other tabular exports/imports as well.
Default Behavior

The annotated format is the default for tabular export/import. To change this behavior, specify freeform in the relevant export/import context.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

- In freeform, the num_rows x num_cols total data entries may be separated with any whitespace including spaces, tabs, and newlines. In this format, vectors may therefore appear as a single row or single column (or mixture; entries will populate the vector in order).

- Some TPLs like SCOLIB and JEGA manage their own file I/O and only support the freeform option.

Examples

Export a freeform tabular file containing only data for variables and responses. Input file fragment:

```environment
tabular_data
tabular_data_file = ‘dakota_summary.dat’
freeform```

Resulting tabular file:

<p>| | | | | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>0.9</td>
<td>1.1</td>
<td>0.0002</td>
<td>0.26</td>
<td>0.76</td>
</tr>
<tr>
<td>0.90009</td>
<td>1.1</td>
<td>0.0001996404857</td>
<td>0.2601620081</td>
<td>0.759955</td>
</tr>
<tr>
<td>0.89991</td>
<td>1.1</td>
<td>0.0002003604863</td>
<td>0.2598380081</td>
<td>0.760045</td>
</tr>
</tbody>
</table>

...
Description

By default, files for tabular data imports are expected to contain columns for all variables, active and inactive. The keyword `active_only` indicates that the file to import contains only the active variables.

This option should only be used in contexts where the inactive variables have no influence, for example, building a surrogate over active variables, with the state variables held at nominal. It should not be used in more complex nested contexts, where the values of inactive variables are relevant to the function evaluations used to build the surrogate.

7.3.3.3 multipoint

- **Keywords Area**
  - model
  - surrogate
  - multipoint

Construct a surrogate from multiple existing training points

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>Multipoint Surrogate (Group 1)</td>
<td>tana</td>
<td>Local multi-point model via two-point nonlinear approximation</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>qmea</td>
<td>Multi-point surrogate approximation based on QMEA algorithm</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>actual_model_pointer</td>
<td>Pointer to specify a &quot;truth&quot; model, from which to construct a surrogate</td>
</tr>
</tbody>
</table>

Description

Multipoint approximations use data from previous design points to improve the accuracy of local approximations. The data often comes from the current and previous iterates of a minimization algorithm.

Currently, only the Two-point Adaptive Nonlinearity Approximation (TANA-3) method of [95] is supported with the `tana` keyword.

The truth model to be used to generate the value/gradient data used in the approximation is identified through the `required actual_model_pointer specification`. 
See Also

These keywords may also be of interest:

- local
- global
- hierarchical

7.3.3.3.1 tana

- Keywords Area
- model
- surrogate
- multipoint
- tana

Local multi-point model via two-point nonlinear approximation

Specification

**Alias:** none

**Argument(s):** none

Description

**TANA** stands for Two Point Adaptive Nonlinearity Approximation.

The TANA-3 method\[95\] is a multipoint approximation method based on the two point exponential approximation\[25\]. This approach involves a Taylor series approximation in intermediate variables where the powers used for the intermediate variables are selected to match information at the current and previous expansion points.

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

Theory

The form of the TANA model is:

\[
\hat{f}(\mathbf{x}) \approx f(\mathbf{x}_2) + \sum_{i=1}^{n} \frac{\partial f}{\partial x_i}(\mathbf{x}_2) \frac{x_i^{1-p_i}}{p_i} (x_i^{p_i} - x_i^{p_i}) + \frac{1}{2} \mathbf{e}(\mathbf{x}) \sum_{i=1}^{n} (x_i^{p_i} - x_i^{p_i})^2
\]

where \(n\) is the number of variables and:

\[
p_i = 1 + \ln \left[ \frac{\frac{\partial f}{\partial x_i}(\mathbf{x}_1)}{\frac{\partial f}{\partial x_i}(\mathbf{x}_2)} \right] \ln \left[ \frac{x_i^{p_i}}{x_i^{p_i}} \right]
\]

\(H\) is determined by

\[
H = 2 \left[ f(\mathbf{x}_1) - f(\mathbf{x}_2) - \sum_{i=1}^{n} \frac{\partial f}{\partial x_i}(\mathbf{x}_2) \frac{x_i^{1-p_i}}{p_i} (x_i^{p_i} - x_i^{p_i}) \right]
\]

and \(\mathbf{x}_2\) and \(\mathbf{x}_1\) are the current and previous expansion points. Prior to the availability of two expansion points, a first-order Taylor series is used.
7.3.3.2 qmea

• Keywords Area
• model
• surrogate
• multipoint
• qmea

Multi-point surrogate approximation based on QMEA algorithm

Specification

Alias: none
Argument(s): none

Description

The Quadratic Multipoint Exponential Approximation (QMEA) builds a multi-point approximation from values and gradients at multiple expansion points. It is a generalization of the two-point exponential approximation (TPEA). This capability is experimental.

7.3.3.3 actual_model_pointer

• Keywords Area
• model
• surrogate
• multipoint
• actual_model_pointer

Pointer to specify a "truth" model, from which to construct a surrogate

Topics

This keyword is related to the topics:

• block_pointer

Specification

Alias: truth_model_pointer
Argument(s): STRING

Description

This must point to a model block, identified by id_model. That model will be run to generate training data, from which a surrogate model will be constructed.

See block_pointer for details about pointers.
Build a locally accurate surrogate from data at a single point

## Specification

**Alias:** none

**Argument(s):** none

### Child Keywords:

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<td><strong>Required</strong></td>
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<td>taylor_series</td>
<td>Construct a Taylor Series expansion around a point</td>
</tr>
<tr>
<td><strong>Required</strong></td>
<td></td>
<td>actual_model_pointer</td>
<td>Pointer to specify a “truth” model, from which to construct a surrogate</td>
</tr>
</tbody>
</table>

### Description

Local approximations use value, gradient, and possibly Hessian data from a single point to form a series expansion for approximating data in the vicinity of this point.

The currently available local approximation is the `taylor_series` selection.

The truth model to be used to generate the value/gradient/Hessian data used in the series expansion is identified through the required `actual_model_pointer` specification. The use of a model pointer (as opposed to an interface pointer) allows additional flexibility in defining the approximation. In particular, the derivative specification for the truth model may differ from the derivative specification for the approximation, and the truth model results being approximated may involve a model recursion (e.g., the values/gradients from a nested model).

### See Also

These keywords may also be of interest:

- global
- hierarchical
- multipoint

### 7.3.3.4.1 taylor_series

- Keywords Area
- model
- surrogate
Construct a Taylor Series expansion around a point

**Specification**

*Alias:* none  
*Argument(s):* none

**Description**

The Taylor series model is purely a local approximation method. That is, it provides local trends in the vicinity of a single point in parameter space.

The order of the Taylor series may be either first-order or second-order, which is automatically determined from the gradient and Hessian specifications in the responses specification (see responses for info on how to specify gradients and Hessians) for the truth model.

**Known Issue:** When using discrete variables, there have been sometimes significant differences in surrogate behavior observed across computing platforms in some cases. The cause has not yet been fully diagnosed and is currently under investigation. In addition, guidance on appropriate construction and use of surrogates with discrete variables is under development. In the meantime, users should therefore be aware that there is a risk of inaccurate results when using surrogates with discrete variables.

**Theory**

The first-order Taylor series expansion is:

\[
\hat{f}(x) \approx f(x_0) + \nabla_x f(x_0)^T (x - x_0)
\]  

and the second-order expansion is:

\[
\hat{f}(x) \approx f(x_0) + \nabla_x f(x_0)^T (x - x_0) + \frac{1}{2} (x - x_0)^T \nabla^2_x f(x_0) (x - x_0)
\]

where \(x_0\) is the expansion point in \(n\)-dimensional parameter space and \(f(x_0), \nabla_x f(x_0),\) and \(\nabla^2_x f(x_0)\) are the computed response value, gradient, and Hessian at the expansion point, respectively.

As dictated by the responses specification used in building the local surrogate, the gradient may be analytic or numerical and the Hessian may be analytic, numerical, or based on quasi-Newton secant updates.

In general, the Taylor series model is accurate only in the region of parameter space that is close to \(x_0\). While the accuracy is limited, the first-order Taylor series model reproduces the correct value and gradient at the point \(x_0\), and the second-order Taylor series model reproduces the correct value, gradient, and Hessian. This consistency is useful in provably-convergent surrogate-based optimization. The other surface fitting methods do not use gradient information directly in their models, and these methods rely on an external correction procedure in order to satisfy the consistency requirements of provably-convergent SBO.

**7.3.3.4.2 actual_model_pointer**

- Keywords Area
- model
- surrogate
- local
**actual_model_pointer**

Pointer to specify a "truth" model, from which to construct a surrogate

**Topics**

This keyword is related to the topics:

- block_pointer

**Specification**

**Alias:** truth_model_pointer

**Argument(s):** STRING

**Description**

This must point to a model block, identified by id_model. That model will be run to generate training data, from which a surrogate model will be constructed.

See block_pointer for details about pointers.

**7.3.3.5 hierarchical**

- **Keywords Area**
- model
- surrogate
- hierarchical

Hierarchical surrogates use an ordered hierarchy of model fidelities and/or resolutions to trade accuracy versus cost.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<tr>
<td>Required</td>
<td></td>
<td>ordered_model_ -fidelities</td>
<td>Specification of an hierarchy of model fidelities, ordered from low to high.</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>correction</td>
<td>Correction approaches for surrogate models</td>
</tr>
</tbody>
</table>
Description

Multifidelity modeling involves the use of lower-fidelity physics-based models as surrogates for a high-fidelity "truth" model. These low-fidelity models can involve variations in model form, resolution, or both. Model form variations for reduced fidelity may involve omitted physics or solution of approximated or averaged physics. Resolution variations typically involve coarsened space-time discretizations (e.g., h- derefinement), reduced element order (e.g., p- derefinement), or loosened convergence tolerances. Unlike local, global, and multipoint surrogates, these approximations are not data-driven (no high-fidelity data required for construction) and are stand-alone simulation models in their own right. As an example from computational fluid dynamics (CFD), both model form and resolution are varied when an inviscid, incompressible Euler model on a coarse discretization is used as a low-fidelity surrogate for a high-fidelity large eddy simulation model on a fine discretization.

The required ordered_model_fidelities specification points to a sequence of model specifications of varying fidelity, ordered from lowest to highest fidelity. The highest fidelity model provides the "truth" model, and each of the lower fidelity alternatives provides different levels of approximation at different levels of cost. This specification defines the sequence of model forms, and each model specification identified in this ordered listing may identify a set of resolution controls. Either or both of these sequences may be specified, and this is referred to as a "multilevel hierarchy" in the case of a hierarchy of resolutions (one entry in ordered_model_fidelities that includes active resolution control), a "multifidelity hierarchy" in the case of a hierarchy of model forms (multiple entries in ordered_model_fidelities without active resolution control), or a "multilevel-multifidelity hierarchy" in the case of two-dimensional hierarchy including both model forms and resolutions (multiple entries in ordered_model_fidelities with one or more including active resolution control). Note that the multilevel-multifidelity case can be "ragged" in the sense that not all models need to provide the same number of (or any) resolution controls.

The correction specification identifies what type (additive, multiplicative, combined) and order (zeroth_order, first_order, second_order) of correction technique will be applied to the low fidelity results in order to match high fidelity results (value and potentially gradient and Hessian) at one or more points. As described below, this is essential in the optimization context.

Use cases

In multifidelity surrogate-based optimization (SBO), the search algorithm relies primarily on the lower fidelity models, which are corrected for consistency with higher fidelity models. The higher fidelity models are used primarily for verifying candidate steps based on solution of low fidelity approximate subproblems and for updating low fidelity corrections. In the hierarchical SBO case (as compared to SBO with data fits), the correction specification is required, since the omission of a correction technique would effectively eliminate the purpose of the high fidelity model (to use a low fidelity model without corrections, then a single model can be used rather than a hierarchical model). Refer to global for additional information on available correction approaches.

In multifidelity uncertainty quantification (UQ), response differences are tracked for purposes of decomposing variance across model/resolution levels or for constructing separate discrepancy emulators. In this context, correction specifications are still valid for defining discrepancy emulation details but they are optional with the most common cases used as defaults.

Examples

Theory:

See Also

These keywords may also be of interest:

- global
- local
- multipoint
Keywords Area

- multilevel_sampling
- multilevel_polynomial_chaos
- multifidelity_polynomial_chaos
- multifidelity_stoch_collocation
- multilevel_function_train
- multifidelity_function_train
- surrogate_based_local

7.3.3.5.1 ordered_model_fidelities

- Keywords Area
- model
- surrogate
- hierarchical
- ordered_model_fidelities

Specification of an hierarchy of model fidelities, ordered from low to high.

Specification

**Alias:** model_fidelity_sequence

**Argument(s):** STRINGLIST

Description

A hierarchical surrogate model manages an ordered set of model fidelities, each of which may in turn involve multiple discretization levels (in the case of a simulation model) or additional model recursions.

The ordering is assumed to be from lowest fidelity to highest fidelity, as dictated by an accuracy versus cost trade-off. Corresponding sequence specifications within methods (e.g., quadrature_order_sequence, sparse_grid_level_sequence, expansion_order_sequence, etc. within stochastic expansion methods) should be synchronized with this model order.

**Additional Discussion**

Internal to the model, only one low fidelity model instance and one high fidelity model instance are active at any given time, although various optimization and UQ algorithms can be used to traverse deep multilevel and multifidelity hierarchies by activating different model combinations and different response modes within the hierarchical model infrastructure.

**Examples**

```dakota
model, id_model = 'HIERARCH'
  surrogate hierarchical
  ordered_model_fidelities = 'LF' 'MF' 'HF'
  correction additive zeroth_order

model, id_model = 'LF'
simulation
  interface_pointer = 'LF_DRIVER'
```
model,  
    id_model = 'MF'  
    simulation  
        interface_pointer = 'MF_DRIVER'

model,  
    id_model = 'HF'  
    simulation  
        interface_pointer = 'HF_DRIVER'

See Also

These keywords may also be of interest:

- multilevel_sampling
- polynomial_chaos
- stoch_collocation
- surrogate_based_local

7.3.3.5.2 correction

- Keywords Area
- model
- surrogate
- hierarchical
- correction

Correction approaches for surrogate models

Specification

Alias: none

Argument(s): none

Default: no surrogate correction

Child Keywords:

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<th>Dakota Keyword</th>
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<td>zeroth_order</td>
<td>Specify that truth values must be matched.</td>
<td></td>
</tr>
<tr>
<td></td>
<td>first_order</td>
<td>Specify that truth values and gradients must be matched.</td>
<td></td>
</tr>
</tbody>
</table>
**Description**

Some of the surrogate model types support the use of correction factors that improve the local accuracy of the surrogate models.

The correction specification specifies that the approximation will be corrected to match truth data, either matching truth values in the case of zeroth_order matching, matching truth values and gradients in the case of first_order matching, or matching truth values, gradients, and Hessians in the case of second_order matching. For additive and multiplicative corrections, the correction is local in that the truth data is matched at a single point, typically the center of the approximation region. The additive correction adds a scalar offset (zeroth_order), a linear function (first_order), or a quadratic function (second_order) to the approximation to match the truth data at the point, and the multiplicative correction multiplies the approximation by a scalar (zeroth_order), a linear function (first_order), or a quadratic function (second_order) to match the truth data at the point. The additive first_order case is due to[59] and the multiplicative first_order case is commonly known as beta correction[41]. For the combined correction, the use of both additive and multiplicative corrections allows the satisfaction of an additional matching condition, typically the truth function values at the previous correction point (e.g., the center of the previous trust region). The combined correction is then a multipoint correction, as opposed to the local additive and multiplicative corrections. Each of these correction capabilities is described in detail in[24].

The correction factors force the surrogate models to match the true function values and possibly true function derivatives at the center point of each trust region. Currently, Dakota supports either zeroth-, first-, or second-order accurate correction methods, each of which can be applied using either an additive, multiplicative, or combined correction function. For each of these correction approaches, the correction is applied to the surrogate model and the corrected model is then interfaced with whatever algorithm is being employed. The default behavior is that no correction factor is applied.

The simplest correction approaches are those that enforce consistency in function values between the surrogate and original models at a single point in parameter space through use of a simple scalar offset or scaling applied to the surrogate model. First-order corrections such as the first-order multiplicative correction (also known as beta correction[14]) and the first-order additive correction[59] also enforce consistency in the gradients and provide a much more substantial correction capability that is sufficient for ensuring provable convergence in SBO algorithms. SBO convergence rates can be further accelerated through the use of second-order corrections which also enforce consistency in the Hessians[24], where the second-order information may involve analytic, finite-difference, or quasi-Newton Hessians.

Correcting surrogate models with additive corrections involves

\[
\hat{f}_{hiq}(x) = f_{lo}(x) + \alpha(x)
\]  

(7.9)
where multifidelity notation has been adopted for clarity. For multiplicative approaches, corrections take the form

\[ \hat{f}_{hi} (x) = f_{lo}(x) \beta(x) \]  

(7.10)

where, for local corrections, \( \alpha(x) \) and \( \beta(x) \) are first or second-order Taylor series approximations to the exact correction functions:

\[ \alpha(x) = A(x_c) + \nabla A(x_c)^T (x - x_c) + \frac{1}{2} (x - x_c)^T \nabla^2 A(x_c) (x - x_c) \]  

(7.11)

\[ \beta(x) = B(x_c) + \nabla B(x_c)^T (x - x_c) + \frac{1}{2} (x - x_c)^T \nabla^2 B(x_c) (x - x_c) \]  

(7.12)

where the exact correction functions are

\[ A(x) = f_{hi}(x) - f_{lo}(x) \]  

(7.13)

\[ B(x) = \frac{f_{hi}(x)}{f_{lo}(x)} \]  

(7.14)

Refer to\[24\] for additional details on the derivations.

A combination of additive and multiplicative corrections can provide for additional flexibility in minimizing the impact of the correction away from the trust region center. In other words, both additive and multiplicative corrections can satisfy local consistency, but through the combination, global accuracy can be addressed as well. This involves a convex combination of the additive and multiplicative corrections:

\[ \hat{f}_{hi}(x) = \gamma \hat{f}_{ha}(x) + (1 - \gamma) \hat{f}_{hb}(x) \]

where \( \gamma \) is calculated to satisfy an additional matching condition, such as matching values at the previous design iterate. It should be noted that in both first order correction methods, the function \( \hat{f}(x) \) matches the function value and gradients of \( f_t(x) \) at \( x = x_c \). This property is necessary in proving that the first order-corrected SBO algorithms are provably convergent to a local minimum of \( f_t(x) \). However, the first order correction methods are significantly more expensive than the zeroth order correction methods, since the first order methods require computing both \( \nabla f_t(x_c) \) and \( \nabla f_s(x_c) \). When the SBO strategy is used with either of the zeroth order correction methods, or with no correction method, convergence is not guaranteed to a local minimum of \( f_t(x) \). That is, the SBO strategy becomes a heuristic optimization algorithm. From a mathematical point of view this is undesirable, but as a practical matter, the heuristic variants of SBO are often effective in finding local minima.

**Usage guidelines**

- Both the additive zeroth_order and multiplicative zeroth_order correction methods are “free” since they use values of \( f_t(x_c) \) that are normally computed by the SBO strategy.

- The use of either the additive first_order method or the multiplicative first_order method does not necessarily improve the rate of convergence of the SBO algorithm.

- When using the first order correction methods, the gradient-related response keywords must be modified to allow either analytic or numerical gradients to be computed. This provides the gradient data needed to compute the correction function.

- For many computationally expensive engineering optimization problems, gradients often are too expensive to obtain or are discontinuous (or may not exist at all). In such cases the heuristic SBO algorithm has been an effective approach at identifying optimal designs[35].
Specify that truth values must be matched.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The correction specification specifies that the approximation will be corrected to match truth data. The keyword zeroth_order matching ensures that truth values are matched.

7.3.3.5.2.2 first_order

- **Keywords Area**
  - model
  - surrogate
  - hierarchical
  - correction
  - first_order

Specify that truth values and gradients must be matched.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

This correction specification specifies that the approximation will be corrected to match truth data. The keyword first_order matching ensures that truth values and gradients are matched.

7.3.3.5.2.3 second_order

- **Keywords Area**
  - model
  - surrogate
  - hierarchical
Specify that truth values, gradients and Hessians must be matched.

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

The correction specification specifies that the approximation will be corrected to match truth data. The keyword `second_order` matching ensures that truth values, gradients and Hessians are matched.

7.3.3.5.2.4 **additive**

- Keywords Area
- model
- surrogate
- hierarchical
- correction
- additive

Additive correction factor for local surrogate accuracy

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Use an additive correction factor to improve the local accuracy of a surrogate.

7.3.3.5.2.5 **multiplicative**

- Keywords Area
- model
- surrogate
- hierarchical
- correction
- multiplicative

Multiplicative correction factor for local surrogate accuracy.
Specification

Alias: none
Argument(s): none

Description

Use a multiplicative correction factor to improve the local accuracy of a surrogate.

7.3.3.5.2.6 combined

- Keywords Area
- model
- surrogate
- hierarchical
- correction
- combined

Multipoint correction for a hierarchical surrogate

Specification

Alias: none
Argument(s): none

Description

For the combined correction, the use of both additive and multiplicative corrections allows the satisfaction of an additional matching condition, typically the truth function values at the previous correction point (e.g., the center of the previous trust region). The combined correction is then a multipoint correction, as opposed to the local additive and multiplicative corrections.

7.3.3.6 non_hierarchical

- Keywords Area
- model
- surrogate
- non_hierarchical

Non-hierarchical surrogates employ an unordered set of lower-fidelity models to approximate a truth reference model at reduced cost.

Specification

Alias: model_ensemble
Argument(s): none
Child Keywords:
### 7.3 model

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<td>truth_model_pointer</td>
<td>Pointer to specify a &quot;truth&quot; model, from which to construct a surrogate</td>
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<td>Required</td>
<td></td>
<td>unordered_model_fidelities</td>
<td>Specification of an unordered ensemble of low-fidelity approximations</td>
</tr>
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</table>

#### See Also

These keywords may also be of interest:

- hierarchical

#### 7.3.3.6.1 truth_model_pointer

- Keywords Area
- model
- surrogate
- non_hierarchical
- truth_model_pointer

Pointer to specify a "truth" model, from which to construct a surrogate

#### Topics

This keyword is related to the topics:

- block_pointer

#### Specification

**Alias:** actual_model_pointer

**Argument(s):** STRING

**Description**

This must point to a model block, identified by `id_model`. That model will be run to generate training data, from which a surrogate model will be constructed.

See `block_pointer` for details about pointers.

#### 7.3.3.6.2 unordered_model_fidelities

- Keywords Area
- model
• surrogate
• non_hierarchical
• unordered_model_fidelities

Specification of an unordered ensemble of low-fidelity approximations

**Specification**

**Alias:** approximation_models

**Argument(s):** STRINGLIST

**Description**

A non_hierarchical surrogate model manages an unordered set of low-fidelity model approximations, each of which may include hyper-parameter resolution controls (in the case of a simulation model) or additional model recursions.

Any corresponding sequence specifications within methods (e.g., quadrature_order_sequence, sparse_grid_level_sequence, expansion_order_sequence, etc. within stochastic expansion methods) should be synchronized with the order in the model listing.

Internal to the non-hierarchical model, subsets of the model ensemble may be active for any given evaluation, as dictated by the iterative algorithm in use.

**Examples**

```plaintext
model,
   id_model = 'NONHIERARCH'
   surrogate non_hierarchical
   unordered_model_fidelities = 'LF1' 'LF2'
   truth_model_pointer = 'HF'

model,
   id_model = 'LF1'
   simulation
   interface_pointer = 'LF1_DRIVER'
   solution_level_cost = 1.

model,
   id_model = 'LF2'
   simulation
   interface_pointer = 'LF2_DRIVER'
   solution_level_cost = 2.4

model,
   id_model = 'HF'
   simulation
   interface_pointer = 'HF_DRIVER'
   solution_level_cost = 256.
```

**See Also**

These keywords may also be of interest:

• hierarchical
A model whose responses are computed through the use of a sub-iterator

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>optional_interface_pointer</td>
<td>Pointer to interface that provides non-nested responses</td>
</tr>
<tr>
<td>Required</td>
<td></td>
<td>sub_method_pointer</td>
<td>The sub_method_pointer specifies the method block for the sub-iterator</td>
</tr>
</tbody>
</table>

**Description**

Instead of appealing directly to a primary interface, a nested model maps variables to responses by executing a secondary iterator, or a "sub-iterator". In other words, a function evaluation of the primary study consists of a solution of an entire secondary study - potentially many secondary function evaluations.

The sub-iterator in turn operates on a sub-model. The sub-iterator responses may be combined with non-nested contributions from an optional interface specification.

A **sub_method_pointer** must be provided in order to specify the method block describing the sub-iterator. The remainder of the model is specified under that keyword.

A **optional_interface_pointer** points to the interface specification and **optional_interface_responses_pointer** points to a responses specification describing the data to be returned by this interface. This interface is used to provide non-nested data, which is then combined with data from the nested iterator using the primary_response_mapping. (See the discussions for the sub_method_pointer and optional_interface_pointer keywords.)

**See Also**

These keywords may also be of interest:

- single
- surrogate
7.3.4.1 optional_interface_pointer

- Keywords Area
- model
- nested
- optional_interface_pointer

Pointer to interface that provides non-nested responses

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Default: no optional interface

Child Keywords:

<table>
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<th>Required/Optional</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>optional_interface(pointer)</td>
<td>responses block that defines non-nested responses</td>
</tr>
</tbody>
</table>

Description

optional_interface_pointer is used to specify an optional interface (using that interface block's id_interface label) to provide non-nested responses to the nested model.

To generate the complete set of primary responses (response functions, objective functions, and calibration terms) expected by the nested model, primary responses from the optional interface may be combined (by addition) with responses derived from sub-iterator results, or they may be presented unchanged to the nested model. The way that Dakota treats primary responses from these two sources depends on the number of rows of and entries in the primary_response_mapping matrix.

To understand how, it is helpful to consider the following expression. Nested model primary responses $r$ are computed from the column vector of optional interface responses $f$, the primary_response_mapping matrix $W$, and the column vector of results from the sub-iterator $s$.

$$\{r\} = \begin{pmatrix} f_0 \\ 0 \end{pmatrix} + \begin{pmatrix} Ws \\ 0 \end{pmatrix}$$

The number of rows in $f$ is equal to the number of primary responses returned by the optional interface, and in $r$, the number of primary responses expected by the nested model. In this expression, $0$ represents “padding” that may be needed by either the optional interface vector or the sub-iterator vector to make them the same dimension as $r$. One or the other of these quantities may be padded, but not both. In terms of the Dakota input, this means that the number of nested model primary responses must equal the larger of these two quantities: the number of optional interface responses or the number of rows in the primary_response_mapping.
The nested model's secondary responses (nonlinear constraints) are the secondary responses from the optional interface augmented by the secondary responses constructed from sub-iterator results. Unlike primary responses from these two sources, which may be combined by adding, it is not possible presently to combine secondary responses. For this reason, the number of nested model's secondary responses must equal the number of secondary responses provided by the optional interface, plus the number provided by the sub-iterator (i.e. the number of rows in the secondary_response_mapping). Secondary responses from the optional interface are inserted first into the nested model's nonlinear constraints; those from the sub-iterator come after. If the nested model has secondary responses and an optional interface is used, it is necessary to specify a separate response block for the optional interface using the optional_interface_response_pointer. Targets and upper and lower bounds for the nonlinear constraints are taken from the nested model's response block.

Examples

The first example illustrates the use of an optional interface and primary response mapping. No secondary responses are present. Assume that the inner method returns two results (suppose its the mean and standard deviation of a single response), so that the dimensions of primary_response_mapping will be interpreted by Dakota as 1x2, just as they are written here.

The nested model (nested) expects two primary responses (outer_resp block), and the optional interface returns two (opt_resp block).

```plaintext
model
  id_model 'nested'
  nested
    optional_interface_pointer 'opt_intf'
      optional_interface_response_pointer 'opt_resp'
    sub_method_pointer 'inner_method'
      primary_response_mapping 1.0 2.0
    responses_pointer 'outer_resp'
    variables_pointer 'outer_vars'
responses
  id_responses 'opt_resp'
  response_functions 2
    descriptors 'opt1' 'opt2'
    no_gradients
    no_hessians
responses
  id_responses 'outer_resp'
  response_functions 2
    descriptors 'out1' 'out2'
    no_gradients
    no_hessians
```

The first nested model response, out1 will be computed $\text{out1} = \text{opt1} + 1.0 \cdot \text{mean} + 2.0 \cdot \text{stddev}$, and the second response out2 will be equal to simply opt2. In the matrix form introduced above, including the implicit padding:

$$\begin{pmatrix}
  \text{out1} \\
  \text{out2}
\end{pmatrix}
= \begin{pmatrix}
  \text{opt1} \\
  \text{opt2}
\end{pmatrix}
+ \begin{pmatrix}
  1.0 \cdot \text{mean} + 2.0 \cdot \text{stddev} \\
  0
\end{pmatrix}
$$

The optional_interface_response_pointer was not strictly needed. Because the number of responses returned by the optional interface and expected by the nested model is the same (and no secondary responses are present), the nested model's response specification alone would have sufficed.

If it were desired to combine opt2 with the sub-iterator's results and set out1 to opt1 with no combination, then the primary_response_mapping matrix could be changed to:

```
primary_response_mapping 0.0 0.0
                        1.0 2.0
```

Secondary responses will be demonstrated in the second example.
The nested model expects a total of two constraints. Because the secondary_response_mapping has one row, one of these constraints will be supplied from the sub-iterator. The other is provided by the optional interface. Secondary responses from the optional iterator come first, and so optc is placed in outc1. outc2 takes the value computed from the sub-iterator results.

### optional_interface_responses_pointer

- **Keywords Area**
  - model
  - nested
  - optional_interface_pointer
  - optional_interface_responses_pointer

Pointer to responses block that defines non-nested responses

### Topics

This keyword is related to the topics:

- **block_pointer**

### Specification

- **Alias:** none
- **Argument(s):** STRING
- **Default:** reuse of top-level responses specification
optional_interface_responses_pointer points to the responses block (specifically, its id_responses label) that defines the non-nested response to return to the nested model. The primary_response_mapping keywords control how these non-nested responses are combined with responses from the nested sub-iterator. See the entry for optional_interface_pointer for a full description. If optional_interface_responses_pointer is not provided, the top-level responses specification is reused.

7.3.4.2 sub_method_pointer

- Keywords Area
- model
- nested
- sub_method_pointer

The sub_method_pointer specifies the method block for the sub-iterator

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>iterator_servers</td>
<td></td>
<td>Specify the number of iterator servers when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>iterator_scheduling</td>
<td></td>
<td>Specify the scheduling of concurrent iterators when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>processors_per_iterator</td>
<td></td>
<td>Specify the number of processors per iterator server when Dakota is run in parallel</td>
</tr>
</tbody>
</table>
The sub_method_pointer specifies the method block for the sub-iterator.

See block_pointer for details about pointers.

A nested model performs an evaluation (variables-to-responses mapping) by running a complete sub-iterator (i.e. method) to completion. The sub-iterator, which is specified using the sub_method_pointer, has its own model. This model, referred to here as the sub-model, possesses its own variables and responses. Prior to every execution of the sub-iterator, information about the nested model's variables is injected into the sub-model's variables. After the sub-iterator has completed, its results are passed back up to the nested model as responses.

Communication of variables information from a nested model to its sub-iterator, and in the opposite direction of sub-iterator results to nested model responses, is called mapping. (This mapping must not be confused with the variables-to-responses mapping that constitutes an evaluation.) Dakota allows considerable power and flexibility in how nested model mappings are performed. They are specified using four keywords: for the variables, primary_variable_mapping and secondary_variable_mapping; and for the responses, primary_response_mapping and secondary_response_mapping. They are described below.

Variable Mappings

In the variable mapping case, primary and secondary variable mapping specifications are used to map from the top-level variables into the sub-model variables. These mappings support three possibilities in any combination: (1) insertion of an active top-level variable value into an identified sub-model distribution parameter for an identified active sub-model variable, (2) insertion of an active top-level variable value into an identified active sub-model variable value, and (3) addition of an active top-level variable value as an inactive sub-model variable, augmenting the active sub-model variables.

For the variable mappings, the primary and secondary specifications are lists of strings that are used to target specific sub-model variables and their sub-parameters, respectively. The primary strings are matched to sub-model variable descriptors such as ‘cdv_1’ (either user-supplied or default labels). The secondary strings are matched to random variable distribution parameters such as ‘mean’ or ‘num_trials’ or design/state variable sub-parameters such as ‘lower_bound’ or ‘upper_bound’.
An important limitation is that real-valued top-level variables must map to real-valued sub-model variables or real-valued sub-parameters, and integer-valued top-level variables must map to either integer-valued sub-model variables or integer-valued sub-parameters. However, as long as these real versus integer constraints are satisfied, mappings are free to cross variable types (design, aleatory uncertain, epistemic uncertain, state) and domain types (continuous, discrete).

Both primary_variable_mapping and secondary_variable_mapping specifications are optional, which is designed to support the following three possibilities:

1. If both primary and secondary variable mappings are specified, then an active top-level variable value will be inserted into the identified sub-parameter (the secondary mapping) for the identified sub-model variable (the primary mapping).

2. If a primary mapping is specified but a secondary mapping is not, then an active top-level variable value will be inserted into the identified sub-model variable value (the primary mapping).

3. If a primary mapping is not specified (corresponding secondary mappings, if specified, are ignored), then an active top-level variable value will be inserted into a corresponding sub-model variable, based on matching of variable types (e.g., top-level and sub-model variable specifications both allocate a set of 'continuous_design' variables which are active at the top level). Multiple sub-model variable types may be updated in this manner, provided that they are all active in the top-level variables. Since there is a direct variable correspondence for these default insertions, sub-model bounds and descriptors are also updated from the top-level bounds and labels in order to eliminate the need for redundant input file specifications. Thus, it is typical for the sub-model variables specification to only contain the minimal required information, such as the number of variables of each type, for these insertion targets. The sub-model must allocate enough space for each of the types that will accept default insertions, and the leading set of matching sub-model variables are updated (i.e., the sub-model may allocate more than needed and the trailing set will be unmodified).

These different variable mapping possibilities may be used in any combination by employing empty strings (") for particular omitted mappings (the number of strings in user-supplied primary and secondary variable mapping specifications must equal the total number of active top-level variables, including both continuous and discrete types). The ordering of the active variables is the same as shown in dakota.input.summary on Input Spec Summary and as presented in the variables section of this manual.

Inactive nested model variables are treated differently from those in the active view. If inactive variables are present at the outer level, then the default type 3 mapping is used for these variables; that is, outer loop inactive variables are inserted into inner loop variables (active or inactive) based on matching of variable types, top-level bounds and labels are also propagated, the inner loop must allocate sufficient space to receive the outer loop values, and the leading subset within this inner loop allocation is updated. This capability is important for allowing nesting beyond two levels, since an active variable at the outer-most loop may become inactive at the next lower level, but still needs to be further propagated down to lower levels in the recursion.

Response Mappings

For the response mappings, the primary and secondary specifications determine how results from the completed sub-iterator are mapped into nested model responses. The response mapping defines a matrix which scales and combines the results from the inner loop into outer loop responses. Each row of the mapping corresponds to one outer loop response, and each column of the mapping corresponds to a result from the inner loop. The results returned from the sub-model are best thought of as forming a column vector; the nested model responses are then the dot-product of the mapping matrix and results vector. The number and type of results that are available and must be accounted for in the response mapping depends on the sub-iterator type:

- optimization: the final objective function(s) and nonlinear constraints
- nonlinear least squares: the final least squares terms and nonlinear constraints
- aleatory uncertainty quantification (UQ): for each response function, a mean statistic, a standard deviation statistic, and all probability/reliability/generalized reliability/response level results for any user-specified response_-levels, probability_levels, reliability_levels, and/or gen_reliability_levels, in that order.
- epistemic and mixed aleatory/epistemic UQ using interval estimation methods: lower and upper interval bounds for each response function.
epistemic and mixed aleatory/epistemic UQ using evidence methods: for each response function, lower and upper interval bounds (belief and plausibility) for all probability/reliability/generalized reliability/response level results computed from any user-specified response_levels, probability_levels, reliability_levels, and/or gen_reliability_levels, in that order. Parameter studies and design of experiments: for optimization and least squares response data sets, the best solution found (lowest constraint violation if infeasible, lowest composite objective function if feasible). For generic response data sets, a best solution metric is not defined, so the sub-iterator response vector is empty in this case.

The primary_response_mapping matrix maps sub-iterator results into top-level objective functions, least squares terms, or generic response functions, depending on the declared top-level response set. The secondary_response_mapping matrix maps sub-iterator results into top-level nonlinear inequality and equality constraints. Alternately, if all the responses of the sub-iterator are to be mapped one-to-one to the top-level nested model responses identity_response_mapping may be specified instead.

Summary

The nested model constructs admit a wide variety of multi-iterator, multi-model solution approaches. For example, optimization within optimization (for hierarchical multidisciplinary optimization), uncertainty quantification within uncertainty quantification (for second-order probability), uncertainty quantification within optimization (for optimization under uncertainty), and optimization within uncertainty quantification (for uncertainty of optima) are all supported, with and without surrogate model indirection. Several examples of nested model usage are provided in the Users Manual, most notably mixed epistemic-aleatory UQ, optimization under uncertainty (OUU), and surrogate-based UQ.

Examples

Two examples are provided to illustrate nested models. The first is a relatively simple case. Although it is somewhat contrived, it demonstrates several features of nested models. A step-by-step explanation is provided below.

```
environment
  method_pointer 'opt'

method
  id_method 'opt'
  async_pattern_search
  model_pointer 'outer_model'
  output verbose

model
  id_model 'outer_model'
  variables_pointer 'outer_vars'
  responses_pointer 'outer_resps'
  nested
    sub_method_pointer 'UQ_method'
      primary_variable_mapping 'x1' 'x2'
      secondary_variable_mapping 'mean' 'mean'
      primary_response_mapping 0 0 0.3 0 0 0.7

variables
  id_variables 'outer_vars'
  continuous_design 2
  descriptors 'x1_mean' 'x2_mean'
  lower_bounds -2.0 -2.0
  upper_bounds 2.0 2.0

responses
  id_responses 'outer_resps'
  objective_functions 1
  descriptors 'sum_p'
  no_gradients
  no_hessians

method
  id_method 'UQ_method'
  sampling samples 30
  seed 1234
  model_pointer 'UQ_model'
```
The example input is an ‘optimization under uncertainty’ or OUU. In an OUU, some statistic of the simulation response is optimized. In this case, a weighted sum of the 90th percentiles of two simulation responses from the text_book driver is being minimized. The uncertainty in these responses is driven by uncertainty in the input variables, which are normally distributed. The means of the uncertain variables are the design variables in the optimization.

In Dakota, this is accomplished by nesting an uncertainty quantification method (sampling), which computes the 90th percentiles, within an optimizer (method-asynch_pattern_search), which iteratively adjusts the variable means to minimize the objective. For every evaluation requested by the optimizer, the design variables (x1_mean and x2_mean in the example) are inserted into the means of the uncertain variables (x1 and x2), then the sampling study is run to completion, and finally the resulting statistics for the responses f1 and f2 are mapped into the responses sum_p.

It may be helpful to sketch out the relationships between blocks as indicated by their various pointers in order to understand the nested structure of the study. The top-level or outer method, ‘opt’, has a model named ‘outer_model’. This model refers to variables and responses blocks (‘outer_vars’ and ‘outer_resps’) that are used by the optimizer. The ‘outer_model’ block also refers to a sub-method, ‘UQ_method’ which defines the UQ study and has its own model, which in turn possess variables, an interface, and responses.

Next, note the primary and secondary variable mappings in ‘outer_model’, which specify that the values of the active design variables x1_mean and x2_mean are to be inserted into the means of the uncertain variables x1 and x2.

The response mapping in ‘outer_model’ is a matrix with a single row and six columns. The single row corresponds to the number of responses in the outer method. As described above, for each of our two sub-model responses, the sampling method returns a mean, a standard deviation, and a single probability level in that order to the nested model. There are a total of six results and accordingly six columns in the mapping matrix. The coefficients 0.3 and 0.7 in the matrix result in a weighted sum of the 90th percentiles:

\[ \text{sum}_p = 0.3 \times (90\text{th percentile of } f1) + 0.7 \times (90\text{th percentile of } f2) \]

The next example is a fragment of an input file, showing only the nested model itself. It illustrates more complex variable and response mappings.
The variable mappings correspond to 4 top-level variables, the first two of which employ the default mappings from active top-level variables to sub-model variables of the same type (option 3 above) and the latter two of which are inserted into the mean distribution parameters of sub-model variables ‘X’ and ‘Y’ (option 1 above).

The response mappings define a 3 by 9 matrix corresponding to 9 inner loop results and 3 outer loop response functions (one primary response function and 2 secondary functions, such as one objective and two constraints). Each row of the response mapping is a vector which is multiplied (i.e., with a dot-product) against the 9 sub-iterator results to determine the outer loop function.

Consider again a UQ example with 3 response functions, each providing a mean, a standard deviation, and one level mapping (if no level mappings are specified, the responses would only have a mean and standard deviation). The primary response mapping can be seen to extract the first result from the inner loop, which would correspond to the mean of the first response function. This mapped sub-iterator response becomes a single objective function, least squares term, or generic response function at the outer level, as dictated by the top-level response specification. The secondary response mapping maps the fourth sub-iterator result plus 3 times the fifth sub-iterator result (mean plus 3 standard deviations) into one top-level nonlinear constraint and the seventh sub-iterator result plus 3 times the eighth sub-iterator result (mean plus 3 standard deviations) into another top-level nonlinear constraint, where these top-level nonlinear constraints may be inequality or equality, as dictated by the top-level response specification.

Note that a common case is for each sub-iterator result to be mapped to a unique outer loop response (for example, in the nested UQ case where one wants to determine an interval on each inner loop statistic). In these simple cases, the response mapping would define an identity matrix and identity_response_mapping may instead be specified.

7.3.4.2.1 iterator_servers

- Keywords Area
- model
- nested
- sub_method_pointer
- iterator_servers

Specify the number of iterator servers when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional iterator_servers specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterations. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

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7.3.4.2.2 iterator_scheduling

• Keywords Area
• model
• nested
• sub_method_pointer
• iterator_scheduling

Specify the scheduling of concurrent iterators when Dakota is run in parallel

Topics

This keyword is related to the topics:

• concurrency_and_parallelism

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Server Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>master</td>
<td>Specify a dedicated master partition for parallel iterator scheduling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel iterator scheduling</td>
</tr>
</tbody>
</table>

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional iterator_scheduling specification supports user override of the automatic parallel configuration for the number of iterator servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.3.4.2.2.1 master

• Keywords Area
• model
• nested
• sub_method_pointer
• iterator_scheduling
• master

Specify a dedicated master partition for parallel iterator scheduling
Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

**Alias:** none

**Argument(s):** none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the iterator servers. This reduces the number of processors available to create servers by 1.

7.3.4.2.2 peer

- Keywords Area
- model
- nested
- sub_method_pointer
- iterator_scheduling
- peer

Specify a peer partition for parallel iterator scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

**Alias:** none

**Argument(s):** none

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to iterator servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.
7.3.4.2.3 processors_per_iterator

• Keywords Area
• model
• nested
• sub_method_pointer
• processors_per_iterator

Specify the number of processors per iterator server when Dakota is run in parallel

Topics

This keyword is related to the topics:

• concurrency_and_parallelism

Specification

Alias: none
Argument(s): INTEGER

Description

An important feature for component-based iterators is that execution of sub-iterator runs may be performed concurrently. The optional processors_per_iterator specification supports user override of the automatic parallel configuration for the number of processors in each iterator server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the iterator parallelism level. Currently, hybrid, multi_start, and pareto_set component-based iterators support concurrency in their sub-iterators. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual[5] for additional information.

7.3.4.2.4 primary_variable_mapping

• Keywords Area
• model
• nested
• sub_method_pointer
• primary_variable_mapping

Primary mapping of top-level variables to sub-model variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: default variable insertions based on variable type
Description

The primary_variable_mapping, secondary_variable_mapping, primary_response_mapping, and secondary_response_mapping keywords control how top-level variables and responses are mapped to variables and responses in the sub-model. Their usage is explained on the parent keyword (sub_method_pointer) page.

7.3.4.2.5 secondary_variable_mapping

- Keywords Area
- model
- nested
- sub_method_pointer
- secondary_variable_mapping

Secondary mapping of top-level variables to sub-model variables

Specification

Alias: none

Argument(s): STRINGLIST

Default: primary mappings into sub-model variables are value-based

Description

The primary_variable_mapping, secondary_variable_mapping, primary_response_mapping, and secondary_response_mapping keywords control how top-level variables and responses are mapped to variables and responses in the sub-model. Their usage is explained on the parent keyword (sub_method_pointer) page.

7.3.4.2.6 primary_response_mapping

- Keywords Area
- model
- nested
- sub_method_pointer
- primary_response_mapping

Mapping of sub-method results to top-level primary responses

Specification

Alias: none

Argument(s): REALLIST

Default: no sub-iterator contribution to primary functions
Description

Matrix that specifies how sub-method results (statistics, best parameters, etc.) map to top-level nested model primary responses (objectives, calibration terms, or response functions). Its usage is explained in detail on the parent keyword \( \text{sub\_method\_pointer} \) page.

7.3.4.2.7 secondary_response_mapping

- Keywords Area
- model
- nested
- sub_method_pointer
- secondary_response_mapping

Mapping of sub-method results to top-level secondary responses

Specification

Alias: none

Argument(s): \text{REALLIST}

Default: no sub-iterator contribution to secondary functions

Description

Matrix that specifies how sub-method results (statistics, best parameters, etc.) map to top-level nested model secondary responses (nonlinear inequality or equality constraints). Its usage is explained in detail on the parent keyword \( \text{sub\_method\_pointer} \) page.

7.3.4.2.8 identity_response_mapping

- Keywords Area
- model
- nested
- sub_method_pointer
- identity_response_mapping

Identity mapping of sub-method results to top-level responses

Specification

Alias: none

Argument(s): none

Default: no sub-iterator contribution to nested model functions
Description

Specifies that the list of sub-method results (statistics, best parameters, etc.) should map one-to-one to the list of top-level nested model responses.

Usage Tips

To use an identity mapping, the number of top-level nested model responses (primary + secondary responses) must equal the number of sub-method results. If receiving an error about this, set output verbose and re-run the Dakota study to see a list of sub-method results responses.

The identity map may not be used in conjunction with optional_interface_pointer, nor with an explicit primary_response_mapping or secondary_response_mapping.

Examples

model
  id_model 'nested'
  nested
  sub_method_pointer 'aleat'
  identity_response_mapping

7.3.4.2.9 Input Spec Summary

This file is derived automatically from dakota.xml, which is used in the generation of parser system files that are compiled into the Dakota executable. Therefore, these files are the definitive source for input syntax, capability options, and associated data inputs. Refer to the Developers Manual information on how to modify the input specification and propagate the changes through the parsing system.

Key features of the input specification and the associated user input files include:

• In the input specification, required individual specifications simply appear, optional individual and group specifications are enclosed in [], required group specifications are enclosed in (), and either-or relationships are denoted by the | symbol. These symbols only appear in dakota.input.summary; they must not appear in actual user input files.

• Keyword specifications (i.e., environment, method, model, variables, interface, and responses) begin with the keyword possibly preceded by white space (blanks, tabs, and newlines) both in the input specifications and in user input files. For readability, keyword specifications may be spread across several lines. Earlier versions of Dakota (prior to 4.1) required a backslash character (\) at the ends of intermediate lines of a keyword. While such backslashes are still accepted, they are no longer required.

• Some of the keyword components within the input specification indicate that the user must supply INTEGER, REAL, STRING, INTEGERLIST, REALLIST, or STRINGLIST data as part of the specification. In a user input file, the "=" is optional, data in a LIST can be separated by commas or whitespace, and the STRING data are enclosed in single or double quotes (e.g., ‘text_book’ or "text_book").

• In user input files, input is largely order-independent (except for entries in lists of data), case insensitive, and white-space insensitive. Although the order of input shown in the Sample Input Files generally follows the order of options in the input specification, this is not required.

• In user input files, specifications may be abbreviated so long as the abbreviation is unique. For example, the npsol_sqp specification within the method keyword could be abbreviated as npsol, but dot_sqp should not be abbreviated as dot since this would be ambiguous with other DOT method specifications.

• In both the input specification and user input files, comments are preceded by #.

• ALIAS refers to synonymous keywords, which often exist for backwards compatibility. Users are encouraged to use the most current keyword.
7.3 model 3527

dakota.input.summary:

KEYWORD01 environment
  [ tabular_data ALIAS tabular_graphics_data
    [ tabular_data_file ALIAS tabular_graphics_file STRING ]
    [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
    | annotated
    | freeform ]
  ]
  [ output_file STRING ]
  [ error_file STRING ]
  [ read_restart STRING
    [ stop_restart INTEGER >= 0 ]
  ]
  [ write_restart STRING ]
  [ output_precision INTEGER >= 0 ]
  [ results_output
    [ results_output_file STRING ]
    [ text ]
    [ hdf5
      [ model_selection
        top_method
        | none
        | all_methods
        | all ]
      [ interface_selection
        none
        | simulation
        | all ]
    ]
  ]
  [ graphics ]
  [ check ]
  [ pre_run
    [ input STRING ]
    [ output STRING
      [ ( custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
      )
      | annotated
      | freeform ]
    ]
  ]
  [ run
    [ input STRING ]
    [ output STRING ]
  ]
  [ post_run
    [ input STRING
      [ ( custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
      )
      | annotated
      | freeform ]
    ]
    [ output STRING ]
  ]
  [ top_method_pointer ALIAS method_pointer STRING ]

KEYWORD method
  [ id_method STRING ]
  [ output
    debug]
<table>
<thead>
<tr>
<th>verbose</th>
<th>normal</th>
<th>quiet</th>
<th>silent</th>
</tr>
</thead>
</table>

{ final_solutions INTEGER >= 0 }

{ hybrid }
  { sequential ALIAS uncoupled }
    { method_name_list STRINGLIST
      [ model_pointer_list STRINGLIST ] }
    { method_pointer_list STRINGLIST
      [ iterator_servers INTEGER > 0 ]
      [ iterator_scheduling
        master
        | peer
      ]
    }
    { processors_per_iterator INTEGER > 0 }
  
  { embedded ALIAS coupled }
    { global_method_name STRING
      [ global_model_pointer STRING ] }
    { local_method_name STRING
      [ local_model_pointer STRING ] }
    { local_method_pointer STRING
      [ local_search_probability REAL ]
      [ iterator_servers INTEGER > 0 ]
      [ iterator_scheduling
        master
        | peer
      ]
    }
    { processors_per_iterator INTEGER > 0 }
  
  { collaborative }
    { method_name_list STRINGLIST
      [ model_pointer_list STRINGLIST ] }
    { method_pointer_list STRINGLIST
      [ iterator_servers INTEGER > 0 ]
      [ iterator_scheduling
        master
        | peer
      ]
    }
    { processors_per_iterator INTEGER > 0 }
  
  { multi_start }
    { method_name STRING
      [ model_pointer STRING ] }
    { method_pointer STRING
      [ random_starts INTEGER
        [ seed INTEGER ]
      ]
    }
    { starting_points REALLIST }
    { iterator_servers INTEGER > 0 }
    { iterator_scheduling
      master
      | peer
    }
    { processors_per_iterator INTEGER > 0 }
  
  { pareto_set }
    { method_name ALIAS opt_method_name STRING
      [ model_pointer ALIAS opt_model_pointer STRING ]
    }
| method_pointer ALIAS opt_method_pointer STRING |
| random_weight_sets INTEGER |
| seed INTEGER |
| weight_sets ALIAS multi_objective_weight_sets REALLIST |
| iterator_servers INTEGER > 0 |
| iterator_scheduling |
| master |
| peer |
| processors_per_iterator INTEGER > 0 |
|
| ( branch_and_bound |
| method_pointer STRING |
| ( method_name STRING |
| model_pointer STRING ) |
| ( scaling ) |
|
| ( surrogate_based_local |
| method_pointer ALIAS approx_method_pointer STRING |
| method_name ALIAS approx_method_name STRING |
| model_pointer ALIAS approx_model_pointer STRING |
| soft_convergence_limit INTEGER |
| truth_surrogate_bypass |
| approx_subproblem |
| original_primary |
| single_objective |
| augmented_lagrangian_objective |
| lagrangian_objective |
| original_constraints |
| linearized_constraints |
| no_constraints ) |
| ( merit_function |
| penalty_merit |
| adaptive_penalty_merit |
| lagrangian_merit |
| augmented_lagrangian_merit ) |
| ( acceptance_logic |
| tr_ratio |
| filter ) |
| ( constraint_relax |
| homotopy ) |
| ( trust_region |
| initial_size REALLIST |
| minimum_size REAL |
| contract_threshold REAL |
| expand_threshold REAL |
| contraction_factor REAL |
| expansion_factor REAL |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| constraint_tolerance REAL |
|
| ( surrogate_based_global |
| method_pointer ALIAS approx_method_pointer STRING |
| method_name ALIAS approx_method_name STRING |
| model_pointer ALIAS approx_model_pointer STRING |
| replace_points |
| max_iterations INTEGER >= 0 |
|
| ( dot_frcg |
| max_iterations INTEGER >= 0 |
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
( model_pointer STRING ]
)

| ( dot_mmfd
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( dot_bfgs
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( dot_slp
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( dot_sqp
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( conmin_frcg
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( conmin_mfd
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ constraint_tolerance REAL ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| ( dl_solver STRING
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)
7.3 model

( npsol_sqp
  [ verify_level INTEGER ]
  [ function_precision REAL ]
  [ linesearch_tolerance REAL ]
  [ convergence_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)

( nlssol_sqp
  [ verify_level INTEGER ]
  [ function_precision REAL ]
  [ linesearch_tolerance REAL ]
  [ convergence_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ constraint_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)

( nlpql_sqp
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)

( optpp_cg
  [ max_step REAL ]
  [ gradient_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)

( optpp_q_newton
  [ search_method
    value_based_line_search
    | gradient_based_line_search
    | trust_region
    | tr_pds
  ]
  [ merit_function
    elbakry
    | argaez_tapia
    | van_shanno
  ]
  [ steplength_to_boundary REAL ]
  [ centering_parameter REAL ]
  [ max_step REAL ]
  [ gradient_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
)

( optpp_fd_newton
  [ search_method
    value_based_line_search
  ]
)
{ optpp_g_newton
  [ search_method
    value_based_line_search
    | gradient_based_line_search
    | trust_region
    | tr_pds
  ]
  [ merit_function
    el_bakry
    | argaez_tapia
    | van_shanno
  ]
  [ steplength_to_boundary REAL ]
  [ centering_parameter REAL ]
  [ max_step REAL ]
  [ gradient_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
}

{ optpp_newton
  [ search_method
    value_based_line_search
    | gradient_based_line_search
    | trust_region
    | tr_pds
  ]
  [ merit_function
    el_bakry
    | argaez_tapia
    | van_shanno
  ]
  [ steplength_to_boundary REAL ]
  [ centering_parameter REAL ]
  [ max_step REAL ]
  [ gradient_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ speculative ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
}

{ optpp_pds
  [ search_scheme_size INTEGER ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
}
7.3 model

```dakota
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)
|
( demo_tpl
[ max_function_evaluations INTEGER >= 0 ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ variable_tolerance REAL ]
[ solution_target ALIAS solution_accuracy REAL ]
[ options_file STRING ]
)
|
( rol
[ max_iterations INTEGER >= 0 ]
[ variable_tolerance REAL ]
[ gradient_tolerance REAL ]
[ constraint_tolerance REAL ]
[ options_file STRING ]
[ scaling ]
[ model_pointer STRING ]
)
|
( asynch_pattern_search ALIAS coliny_apps
[ initial_delta REAL ]
[ contraction_factor REAL ]
[ variable_tolerance REAL ]
[ solution_target ALIAS solution_accuracy REAL ]
[ synchronization ]
[ blocking | nonblocking ]
[ merit_function ]
[ merit_max ]
[ merit_max_smooth ]
[ merit1 ]
[ merit1_smooth ]
[ merit2 ]
[ merit2_smooth ]
[ merit2_squared ]
)
[ constraint_penalty REAL ]
[ smoothing_factor REAL ]
[ constraint_tolerance REAL ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)
|
( mesh_adaptive_search
[ initial_delta REAL ]
[ variable_tolerance REAL ]
[ function_precision REAL ]
[ seed INTEGER > 0 ]
[ history_file STRING ]
[ display_format STRING ]
[ variable_neighborhood_search REAL ]
[ neighbor_order INTEGER > 0 ]
[ display_all_evaluations ]
[ use_surrogate ]
[ inform_search ]
[ optimize ]
)
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)
|
( nowpac
[ trust_region ]
[ initial_size REALLIST ]
```
[ minimum_size REAL ]
[ contract_threshold REAL ]
[ expand_threshold REAL ]
[ contraction_factor REAL ]
[ expansion_factor REAL ]
]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| snowpac
[ seed INTEGER > 0 ]
[ trust_region
[ initial_size REALLIST ]
[ minimum_size REAL ]
[ contract_threshold REAL ]
[ expand_threshold REAL ]
[ contraction_factor REAL ]
[ expansion_factor REAL ]
]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)

| moga
[ fitness_type
layer_rank
| domination_count
]
[ replacement_type
elitist
| roulette_wheel
| unique_roulette_wheel
|
( below_limit REAL
[ shrinkage_fraction ALIAS shrinkage_percentage REAL ]
)
)
[ niching_type
radial REALLIST
|
( max_designs REALLIST
[ num_designs INTEGER >= 2 ]
)
]
[ convergence_type
metric_tracker
[ percent_change REAL ]
[ num_generations INTEGER >= 0 ]
]
[ postprocessor_type
orthogonal_distance REALLIST
]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ population_size INTEGER >= 0 ]
[ log_file STRING ]
[ print_each_pop ]
[ initialization_type
simple_random
| unique_random
| flat_file STRING
]
[ crossover_type
multi_point_binary INTEGER
| multi_point_parameterized_binary INTEGER
| multi_point_real INTEGER
]
7.3 model

```
| ( shuffle_random
|    [ num_parents INTEGER > 0 ]
|    [ num_offspring INTEGER > 0 ]
| )
| [ crossover_rate REAL ]
| )
| [ mutation_type
bit_random |
| replace_uniform |
| ( offset_normal
|    [ mutation_scale REAL ]
| )
| ( offset_cauchy
|    [ mutation_scale REAL ]
| )
| ( offset_uniform
|    [ mutation_scale REAL ]
| )
| [ mutation_rate REAL ]
| )
| [ seed INTEGER > 0 ]
| [ convergence_tolerance REAL ]
| [ model_pointer STRING ]
| )
| ( soga
|    [ fitness_type
merit_function
|    [ constraint_penalty REAL ]
| )
| [ replacement_type
elitist |
favor_feasible |
roulette_wheel |
unique_roulette_wheel 
| )
| [ convergence_type
( best_fitness_tracker
    [ percent_change REAL ]
    [ num_generations INTEGER >= 0 ]
 )
| ( average_fitness_tracker
    [ percent_change REAL ]
    [ num_generations INTEGER >= 0 ]
 )
| [ max_iterations INTEGER >= 0 ]
| [ max_function_evaluations INTEGER >= 0 ]
| [ scaling ]
| [ population_size INTEGER >= 0 ]
| [ log_file STRING ]
| [ print_each_pop ]
| [ initialization_type
simple_random |
| unique_random |
| flat_file STRING 
| )
| [ crossover_type
multi_point_binary INTEGER |
| multi_point_parameterized_binary INTEGER |
| multi_point_real INTEGER 
| ( shuffle_random
|    [ num_parents INTEGER > 0 ]
|    [ num_offspring INTEGER > 0 ]
| )
| [ crossover_rate REAL ]
| )
```
[ mutation_type
  bit_random
  | replace_uniform
  |
  ( offset_normal
    [ mutation_scale REAL ]
  )
  |
  ( offset_cauchy
    [ mutation_scale REAL ]
  )
  |
  ( offset_uniform
    [ mutation_scale REAL ]
  )
  [ mutation_rate REAL ]
]
[ seed INTEGER > 0 ]
[ convergence_tolerance REAL ]
[ model_pointer STRING ]
]
[
  coliny_pattern_search
  [ constant_penalty ]
  [ no_expansion ]
  [ expand_after_success INTEGER ]
  [ pattern_basis
    coordinate
    | simplex
  ]
  [ stochastic ]
  [ total_pattern_size INTEGER ]
  [ exploratory_moves
    multi_step
    | adaptive_pattern
    | basic_pattern
  ]
  [ synchronization
    blocking
    | nonblocking
  ]
  [ contraction_factor REAL ]
  [ constraint_penalty REAL ]
  [ initial_delta REAL ]
  [ variable_tolerance REAL ]
  [ solution_target ALIAS solution_accuracy REAL ]
  [ seed INTEGER > 0 ]
  [ show_misc_options ]
  [ misc_options STRINGLIST ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
  [ model_pointer STRING ]
]
[
  coliny_solis_wets
  [ contract_after_failure INTEGER ]
  [ no_expansion ]
  [ expand_after_success INTEGER ]
  [ constant_penalty ]
  [ contraction_factor REAL ]
  [ constraint_penalty REAL ]
  [ initial_delta REAL ]
  [ variable_tolerance REAL ]
  [ solution_target ALIAS solution_accuracy REAL ]
  [ seed INTEGER > 0 ]
  [ show_misc_options ]
  [ misc_options STRINGLIST ]
  [ max_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ max_function_evaluations INTEGER >= 0 ]
  [ scaling ]
[ model_pointer STRING ]
)
|
| ( coliny_cobyla
| [ initial_delta REAL ]
| [ variable_tolerance REAL ]
| [ solution_target ALIAS solution_accuracy REAL ]
| [ seed INTEGER > 0 ]
| [ show_misc_options ]
| [ misc_options STRINGLIST ]
| [ max_iterations INTEGER >= 0 ]
| [ convergence_tolerance REAL ]
| [ max_function_evaluations INTEGER >= 0 ]
| [ scaling ]
| [ model_pointer STRING ]
)
|
| ( coliny_direct
| [ division
| | major_dimension
| | all_dimensions
| ]
| [ global_balance_parameter REAL ]
| [ local_balance_parameter REAL ]
| [ max_boxsize_limit REAL ]
| [ min_boxsize_limit REAL ]
| [ constraint_penalty REAL ]
| [ solution_target ALIAS solution_accuracy REAL ]
| [ seed INTEGER > 0 ]
| [ show_misc_options ]
| [ misc_options STRINGLIST ]
| [ max_iterations INTEGER >= 0 ]
| [ convergence_tolerance REAL ]
| [ max_function_evaluations INTEGER >= 0 ]
| [ scaling ]
| [ model_pointer STRING ]
)
|
| ( coliny_ea
| [ population_size INTEGER > 0 ]
| [ initialization_type
| | simple_random
| | unique_random
| | flat_file STRING
| ]
| [ fitness_type
| | linear_rank
| | merit_function
| ]
| [ replacement_type
| | random INTEGER
| | chc INTEGER
| | elitist INTEGER
| | [ new_solutions_generated INTEGER ]
| ]
| [ crossover_rate REAL ]
| [ crossover_type
| | two_point
| | blend
| | uniform
| ]
| [ mutation_rate REAL ]
| [ mutation_type
| | replace_uniform
| | [ offset_normal
| | [ mutation_scale REAL ]
| | [ mutation_range INTEGER ]
| ]
| [ offset_cauchy
| [ mutation_scale REAL ]
| [ mutation_range INTEGER ]
]


```
}


coliny_beta

beta_solver_name STRING

[ solution_target ALIAS solution_accuracy REAL ]
[ seed INTEGER > 0 ]
[ show_m misc_options ]
[ misc_options STRINGLIST ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

)

nl2sol

[ function_precision REAL ]
[ absolute_conv_tol REAL ]
[ x_conv_tol REAL ]
[ singular_conv_tol REAL ]
[ singular_radius REAL ]
[ false_conv_tol REAL ]
[ initial_trust_radius REAL ]
[ covariance INTEGER ]
[ regression_diagnostics ]
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ speculative ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

)

nonlinear_cg

[ misc_options STRINGLIST ]
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

)

ncsu_direct

[ solution_target ALIAS solution_accuracy REAL ]
[ min_boxsize_limit REAL ]
[ volume_boxsize_limit REAL ]
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]

)

genie_opt_darts

[ seed INTEGER > 0 ]
[ max_function_evaluations INTEGER >= 0 ]
```
7.3 model

[ scaling ]
[ model_pointer STRING ]
)
|
( genie_direct
[ seed INTEGER > 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ scaling ]
[ model_pointer STRING ]
)
|
( efficient_global
[ initial_samples INTEGER ]
[ seed INTEGER > 0 ]
[ batch_size INTEGER >= 1 ]
[ exploration INTEGER >= 0 ]
[ synchronization
    blocking ]
[ nonblocking ]
)
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ x_conv_tol REAL ]
[ gaussian_process ALIAS kriging
    ( surfpack
    [ export_model
    [ filename_prefix STRING
    [ formats
        [ text_archive ]
        [ binary_archive ]
    ]
    )
    )
    | dakota
    [ experimental
    [ export_model
    [ filename_prefix STRING
    [ formats
        [ text_archive ]
        [ binary_archive ]
    ]
    ]
    [ options_file STRING ]
    ]
][ use_derivatives ]
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
) ]
[ annotated ]
[ freeform ]
[ active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
) ]
[ annotated ]
[ freeform ]
[ max_function ]
[ model_pointer STRING ]
)
|
( surrogate_based_uq
[ samples_on_emulator ALIAS samples INTEGER
[ sample_type ]
[ max_function ]
[ model_pointer STRING ]
)
lhs
| random
|
[rng
  mt19937
  | rnum2
]

[probability_refinement ALIAS sample_refinement
  import
  | adapt_import
  | mm_adapt_import
  [ refinement_samples INTEGERLIST ]
]

[final_moments
  none
  | standard
  | central
]

[response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | reliabilities
    | gen_reliabilities
    [ system
      series
      | parallel
    ]
  ]
]

[probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]]

[reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ]]

[gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]]

[distribution
  cumulative
  | complementary
]

[variance_based_decomp
  [ interaction_order INTEGER > 0 ]
  [ drop_tolerance REAL ]]

[diagonal_covariance
  | full_covariance ]

[import_approx_points_file STRING
  ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform
  [ active_only ]
]

[export_approx_points_file ALIAS export_points_file STRING
  ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform
]

[seed INTEGER > 0 ]
[fixed_seed ]
[model_pointer STRING ]
(function_train
  [ p_refinement
    ( uniform
      increment_start_rank
      | increment_start_order
      | increment_max_rank
      | increment_max_order
      | increment_max_rank_order
    )
  ]
  [ max_refinement_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ metric_scale
    relative
    | absolute
  ]
  [ regression_type
    ls
    | ( rls2
      l2_penalty REAL
    )
  ]
  [ max_solver_iterations INTEGER >= 0 ]
  [ max_cross_iterations INTEGER >= 0 ]
  [ solver_tolerance REAL ]
  [ tensor_grid ]
  collocation_points INTEGER
  | collocation_ratio REAL
  [ rounding_tolerance REAL ]
  [ arithmetic_tolerance REAL ]
  [ start_order ALIAS order INTEGER >= 0
    | dimension_preference REALLIST
  ]
  [ adapt_order ]
  [ kick_order INTEGER > 0 ]
  [ max_order INTEGER >= 0 ]
  [ max_cv_order_candidates INTEGER >= 0 ]
  [ start_rank ALIAS rank INTEGER >= 0
    | adapt_rank
    | kick_rank INTEGER > 0
    | max_rank INTEGER >= 0
    | max_cv_rank_candidates INTEGER >= 0
  ]
  [ samples_on_emulator ALIAS samples INTEGER ]
  [ sample_type
    lhs
    | random
  ]
  [ rng
    mt19937
    | rnum2
  ]
  [ probability_refinement ALIAS sample_refinement
    import
    | adapt_import
    | mm_adapt_import
    [ refinement_samples REALLIST ]
  ]
  [ final_moments
    none
    | standard
    | central
  ]
  [ response_levels REALLIST
    [ num_response_levels INTEGERLIST ]
    [ compute
      probabilities
      | reliabilities
      | gen_reliabilities
      | system
      series
      parallel
    ]]
[ probability_levels REALIST
  [ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALIST
  [ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALIST
  [ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
cumulative
| complementary ]
[ variance_based_decomp
  [ interaction_order INTEGER > 0 ]
  [ drop_tolerance REAL ]
]
[ diagonal_covariance
| full_covariance ]
[ import_approx_points_file STRING
  [ ( custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
  )
| annotated
| freeform ]
| active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
  )
| annotated
| freeform ]
| active_only ]
]
[ multifidelity_function_train
  [ p_refinement
    ( uniform
      increment_start_rank
      | increment_start_order
      | increment_max_rank
      | increment_max_order
      | increment_max_rank_order
    )
  ]
  [ max_refinement_iterations INTEGER >= 0 ]
  [ convergence_tolerance REAL ]
  [ metric_scale
    relative
    | absolute ]
  [ statistics_mode
    active
    | combined ]
  [ allocation_control
    greedy ]
  [ discrepancy_emulation
    distinct ALIAS paired
    | recursive ]
[ rounding_tolerance REAL ]
[ arithmetic_tolerance REAL ]
[ regression_type
  ls
  | ( rls2
  l2_penalty REAL ]
]
[ max_solver_iterations INTEGER >= 0 ]
[ max_cross_iterations INTEGER >= 0 ]
[ solver_tolerance REAL ]
[ response_scaling ]
[ tensor_grid ]
[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
[ collocation_ratio REAL ]
[ start_order_sequence ALIAS order_sequence INTEGERLIST ]
[ dimension_preference REALLIST ]
]
[ adapt_order ]
[ kick_order INTEGER > 0 ]
[ max_order INTEGER >= 0 ]
[ max_cv_order_candidates INTEGER >= 0 ]
[ start_rank_sequence ALIAS rank_sequence INTEGERLIST ]
[ adapt_rank ]
[ kick_rank INTEGER > 0 ]
[ max_rank INTEGER >= 0 ]
[ max_cv_rank_candidates INTEGER >= 0 ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type
  lhs
  | random ]
[ rng
  mt19937
  | rnum2 ]
[ probability_refinement ALIAS sample_refinement import
  | adapt_import
  | mm_adapt_import
  | refinement_samples INTEGERLIST ]
]
[ final_moments
  none
  | standard
  | central ]
[ response_levels REALLIST ]
[ num_response_levels INTEGERLIST ]
[ compute
  probabilities
  | reliabilities
  | gen_reliabilities
  | system
  series
  | parallel ]
]
[ probability_levels REALLIST ]
[ num_probability_levels INTEGERLIST ]
[ reliability_levels REALLIST ]
[ num_reliability_levels INTEGERLIST ]
[ gen_reliability_levels REALLIST ]
[ num_gen_reliability_levels INTEGERLIST ]
[ distribution
  cumulative
  | complementary ]
[ variance_based_decomp
[ interaction_order INTEGER > 0 ]
[ drop_tolerance REAL ]
]
[ diagonal_covariance
[ full_covariance ]
[ import_approx_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform ]
[ active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform ]
]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]
]
{
| multilevel_function_train
[ max_iterations INTEGER >= 0 ]
[ allocation_control
  ( estimator_variance
    | estimator_rate REAL )
] | rank_sampling
]
[ convergence_tolerance REAL ]
[ metric_scale
  relative
  | absolute
]
[ discrepancy_emulation
  distinct ALIAS paired
  | recursive
]
[ rounding_tolerance REAL ]
[ arithmetic_tolerance REAL ]
[ regression_type
  ls
  | ( rls2
    12_penalty REAL
  )
]
[ max_solver_iterations INTEGER >= 0 ]
[ max_cross_iterations INTEGER >= 0 ]
[ solver_tolerance REAL ]
[ response_scaling ]
[ tensor_grid ]
[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
[ collocation_ratio REAL ]
[ start_order_sequence ALIAS order_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
]
[ adapt_order ]
[ kick_order INTEGER > 0 ]
[ max_order INTEGER >= 0 ]
[ max_cv_order_candidates INTEGER >= 0 ]
[ start_rank_sequence ALIAS rank_sequence INTEGERLIST ]
[ adapt_rank ]
[ kick_rank INTEGER > 0 ]
[ max_rank INTEGER >= 0 ]
[ max_cv_rank_candidates INTEGER >= 0 ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type
  lhs |
  random ]
[ rng
  mt19937 |
  rnum2 ]
[ probability_refinement ALIAS sample_refinement
  import |
  adapt_import |
  mm_adapt_import |
  refinement_samples INTEGERLIST ]
[ final_moments
  none |
  standard |
  central ]
[ response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities |
    reliabilities |
    gen_reliabilities |
    system
    series |
    parallel ]
  ]
[ probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]
  ]
[ reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ]
  ]
[ gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]
  ]
[ distribution
  cumulative |
  complementary ]
[ variance_based_decomp
  [ interaction_order INTEGER > 0 ]
  [ drop_tolerance REAL ]
  ]
[ diagonal_covariance |
  full_covariance ]
[ import_approx_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  ) |
  annotated |
  freeform |
  active_only ]
  ]
[ export_approx_points_file ALIAS export_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  ) |
  annotated |
  freeform ]
  ]
[ seed_sequence INTEGERLIST ]
Keywords Area

- fixed_seed
- model_pointer STRING

polynomial_chaos ALIAS nond_polynomial_chaos

p_refinement

- uniform
  - (dimension_adaptive
    - sobol
    - decay
    - generalized)

- max_refinement_iterations INTEGER >= 0
- convergence_tolerance REAL
- metric_scale
  - relative
  - absolute

- quadrature_order INTEGER
  - dimension_preference REALIST
  - nested
  - non_nested

- sparse_grid_level INTEGER
  - dimension_preference REALIST
  - restricted
  - unrestricted
  - nested
  - non_nested

- cubature_integrand INTEGER

- expansion_order INTEGER
  - dimension_preference REALIST
  - basis_type
    - tensor_product
    - total_order
    - (adapted
      - advancements INTEGER
      - soft_convergence_limit INTEGER)

- collocation_points INTEGER
  - (least_squares
    - svd
    - equality_constrained)
  - orthogonal_matching_pursuit ALIAS omp
  - noise_tolerance REALIST
  - basis_pursuit ALIAS bp
  - (basis_pursuit_denoising ALIAS bpdn
    - noise_tolerance REALIST)
  - (least_angle_regression ALIAS lars
    - noise_tolerance REALIST)
  - (least_absolute_shrinkage ALIAS lasso
    - noise_tolerance REALIST
    - l2_penalty REAL)
  - cross_validation
  - noise_only
  - max_cv_order_candidates INTEGER >= 0
7.3 model

[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]

| ( collocation_ratio REAL
| [ { least_squares
| svd
| equality_constrained ]
| )
| ( orthogonal_matching_pursuit ALIAS omp
| [ noise_tolerance REALLIST ]
| ]
| [ basis_pursuit ALIAS bp
| ( basis_pursuit_denoising ALIAS bpdn
| [ noise_tolerance REALLIST ]
| )
| [ least_angle_regression ALIAS lars
| [ noise_tolerance REALLIST ]
| )
| ( least_absolute_shrinkage ALIAS lasso
| [ noise_tolerance REALLIST ]
| [ l2_penalty REAL ]
| ]
| [ cross_validation
| [ noise_only ]
| [ max_cv_order_candidates INTEGER >= 0 ]
| ]
| [ ratio_order REAL ]
| [ response_scaling ]
| [ use_derivatives ]
| [ tensor_grid ]
| [ reuse_points ALIAS reuse_samples ]
| [ max_solver_iterations INTEGER >= 0 ]
|)
| ( expansion_samples INTEGER
| [ reuse_points ALIAS reuse_samples ]
|)
[ import_build_points_file ALIAS import_points_file STRING
[ [ { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
| )
| annotated
| [ freeform ]
| [ active_only ]
| ]
|)
| ( orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
| collocation_points INTEGER
| [ tensor_grid INTEGERLIST ]
| [ reuse_points ALIAS reuse_samples ]
| [ import_build_points_file ALIAS import_points_file STRING
[ [ { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
| )
| annotated
| [ freeform ]
| [ active_only ]
| ]
|)
| import_expansion_file STRING  
| askey                  
| wiener                 
| normalized             
| export_expansion_file STRING  
| samples_on_emulator ALIAS samples INTEGER  
| sample_type lhs        
| random                 
| rng                    
| mt19937                
| rnum2                  
| probability_refinement ALIAS sample_refinement import  
| adapt_import           
| mm_adapt_import        
| refinement_samples INTEGERLIST  
| final_moments none     
| standard               
| central                
| response_levels REALLIST  
| num_response_levels INTEGERLIST  
| compute probabilities  
| reliabilities          
| gen_reliabilities      
| system series          
| parallel               
|                      
| probability_levels REALLIST  
| num_probability_levels INTEGERLIST  
|                      
| reliability_levels REALLIST  
| num_reliability_levels INTEGERLIST  
|                      
| gen_reliability_levels REALLIST  
| num_gen_reliability_levels INTEGERLIST  
|                      
| distribution cumulative  
| complementary         
|                      
| variance_based_decomp  
| interaction_order INTEGER > 0  
| drop_tolerance REAL     
|                      
| diagonal_covariance     
| full_covariance         
|                      
| import_approx_points_file STRING  
| ( custom_annotated  
| header ]           
| eval_id ]          
| interface_id ]     
| annotated          
| freeform ]         
| active_only ]      
|                      
| export_approx_points_file ALIAS export_points_file STRING  
| ( custom_annotated  
| header ]           
| eval_id ]          
| interface_id ]     
| annotated  

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| freeform |
| seed INTEGER > 0 |
| fixed_seed |
| model_pointer STRING |
|
( multifidelity_polynomial_chaos |
| p_refinement |
| uniform |
| ( dimension_adaptive |
| sobol |
| decay |
| generalized |
) |
| max_refinement_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| metric_scale |
| relative |
| absolute |
| statistics_mode |
| active |
| combined |
| allocation_control |
| greedy |
| discrepancy_emulation |
| distinct ALIAS paired |
| recursive |
| ( quadrature_order_sequence INTEGERLIST |
| dimension_preference REALLIST |
| nested |
| non_nested |
) |
| sparse_grid_level_sequence INTEGERLIST |
| dimension_preference REALLIST |
| restricted |
| unrestricted |
| nested |
| non_nested |
) |
| expansion_order_sequence INTEGERLIST |
| dimension_preference REALLIST |
| basis_type |
| tensor_product |
| total_order |
| ( adapted |
| advancements INTEGER |
| soft_convergence_limit INTEGER |
) |
| collocation_ratio REAL |
| collocation_points_sequence ALIAS pilot_samples INTEGERLIST |
| ( least_squares |
| svd |
| equality_constrained |
) |
| ( orthogonal_matching_pursuit ALIAS omp |
| noise_tolerance REALLIST |
) |
| basis_pursuit ALIAS bp |
| ( basis_pursuit_denoising ALIAS bpdn |
| noise_tolerance REALLIST |
(least_angle_regression ALIAS lars
[noise_tolerance REALIST]
)

(least_absolute_shrinkage ALIAS lasso
[noise_tolerance REALIST]
[l2_penalty REAL]
)[cross_validation
[noise_only]
[max_cv_order_candidates INTEGER >= 0]
)[ratio_order REAL]
[response_scaling]
[use_derivatives]
[tensor_grid]
[reuse_points ALIAS reuse_samples]
[max_solver_iterations INTEGER >= 0]
)

(expansion_samples_sequence INTEGERLIST
[reuse_points ALIAS reuse_samples]
)

[import_build_points_file ALIAS import_points_file STRING
[custom_annotated
[header]
[eval_id]
[interface_id]]
)[annotated
[freeform]
[active_only]]
)

(orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
collocation_points_sequence INTEGERLIST
[tensor_grid INTEGERLIST]
[reuse_points ALIAS reuse_samples]
[import_build_points_file ALIAS import_points_file STRING
[custom_annotated
[header]
[eval_id]
[interface_id]]
)[annotated
[freeform]
[active_only]]
)

[askey
[wiener]
[normalized]
[export_expansion_file STRING]
[samples_on_emulator ALIAS samples INTEGER]
[sample_type
[lhs]
[random]]
[rng
[mt19937]
[rnum2]]

(probability_refinement ALIAS sample_refinement
import
[adapt_import
[mm_adapt_import
[refinement_samples INTEGERLIST]]]
)
[final_moments
[none]
7.3 model

[ standard
| central
]
[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
  probabilities
  | reliabilities
  | gen_reliabilities
  | system
  series
  | parallel
 ]
]
[ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
[ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  cumulative
  | complementary
]
[ variance_based_decomp
  [ interaction_order INTEGER > 0 ]
  [ drop_tolerance REAL ]
]
[ diagonal_covariance
| full_covariance ]
[ import_approx_points_file STRING
[ { custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
}
| annotated
| freeform
| active_only
]
[ export_approx_points_file ALIAS export_points_file STRING
[ { custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
}
| annotated
| freeform
]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]
]
[ { multilevel_polynomial_chaos
  [ max_iterations INTEGER >= 0 ]
  [ allocation_control
    { estimator_variance
      [ estimator_rate REAL ]
    } |
    rip_sampling
  ]
  [ convergence_tolerance REAL ]
  [ metric_scale
    relative
    | absolute
  ]
  [ discrepancy_emulation
    distinct ALIAS paired
  ]
]
recursive

(expansion_order_sequence INTEGERLIST
[ dimension_preference REALLIST ]
[ basis_type
tensor_product
| total_order |
| ( adapted
[ advancements INTEGER ]
[ soft_convergence_limit INTEGER ]
)
]
(collocation_ratio REAL
[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
[ { least_squares
| svd
| equality_constrained }
]
| ( orthogonal_matching_pursuit ALIAS omp
[ noise_tolerance REALLIST ]
]
| basis_pursuit ALIAS bp
| ( basis_pursuit_denoising ALIAS bpdn
[ noise_tolerance REALLIST ]
]
| ( least_angle_regression ALIAS lars
[ noise_tolerance REALLIST ]
)
| ( least_absolute_shrinkage ALIAS lasso
[ noise_tolerance REALLIST ]
[ l2_penalty REAL ]
]
| cross_validation
[ noise_only ]
[ max_cv_order_candidates INTEGER >= 0 ]
]
[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]
)
]
(expansion_samples_sequence INTEGERLIST
[ reuse_points ALIAS reuse_samples ]
)
[ import_build_points_file ALIAS import_points_file STRING
[ { custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
]
| annotated
| freeform ]
[ active_only ]
]
)
}
(orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
collocation_points_sequence INTEGERLIST
collocation_points_sequence
[ tensor_grid INTEGERLIST ]
[ reuse_points ALIAS reuse_samples ]
[ import_build_points_file ALIAS import_points_file STRING
[ { custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
]]
7.3 model 3553

)
| annotated
| freeform ]
| active_only ]
|
]
[ askey
| wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type
lhs
| random ]
[ rng
mt19937
| rnum2 ]
[ probability_refinement ALIAS sample_refinement
import
| adapt_import
| mm_adapt_import
[ refinement_samples INTEGERLIST ]
]
[ final_moments
none
| standard
| central ]
[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| reliabilities
| gen_reliabilities
[ system
series
| parallel ]
]
[ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
[ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
cumulative
| complementary ]
[ variance_based_decomp
[ interaction_order INTEGER > 0 ]
| drop_tolerance REAL ]
]
[ diagonal_covariance
| full_covariance ]
[ import_approx_points_file STRING
[ ( custom_annotated
[ header ]
| eval_id ]
| interface_id ]
]
| annotated
| freeform ]
| active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated

[ header ]
[ eval_id ]
[ interface_id ]
[ annotated ]
[ freeform ]
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ model_pointer STRING ]

[ stoch_collocation ALIAS nond_stoch_collocation ]
[ ( p_refinement ]
  [ uniform ]
  [ ( dimension_adaptive ]
    [ sobol ]
    [ generalized ]
  [ ) ]
[ ) ]
[ h_refinement ]
  [ uniform ]
  [ ( dimension_adaptive ]
    [ sobol ]
    [ generalized ]
    [ ) ]
  [ ) ]
[ max_refinement_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ metric_scale ]
  [ relative ]
  [ absolute ]
[ ) ]
[ quadrature_order INTEGER ]
  [ dimension_preference REALLIST ]
  [ nested ]
  [ non_nested ]
[ ) ]
[ sparse_grid_level INTEGER ]
  [ dimension_preference REALLIST ]
  [ nodal ]
  [ hierarchical ]
  [ restricted ]
  [ unrestricted ]
  [ nested ]
  [ non_nested ]
[ ) ]
[ piecewise ]
[ askey ]
[ wiener ]
[ use_derivatives ]
[ samples_on_emulator ALIAS samples INTEGER ]
[ sample_type ]
  [ lhs ]
  [ random ]
[ ) ]
[ rng ]
  mt19937
  rnum2
[ ) ]
[ probability_refinement ALIAS sample_refinement ]
  import ]
  [ adapt_import ]
  [ mm_adapt_import ]
  [ refinement_samples INTEGERLIST ]
[ ) ]
[ final_moments ]
  none
7.3 model 3555

[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
  probabilities
  | reliabilities
  | gen_reliabilities
  | system
series
  | parallel
] ]
}
[ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
[ num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  cumulative
  | complementary
] ]
[ variance_based_decomp
[ interaction_order INTEGER > 0 ]
[ drop_tolerance REAL ]
]
[ diagonal_covariance
| full_covariance ]
[ import_approx_points_file STRING
[ ( custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
  )
| annotated
  | freeform ]
  | active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
  )
| annotated
  | freeform ]
  | active_only ]
  ]
[ seed INTEGER > 0 ]
[ fixed_seed ]
[ model_pointer STRING ]
]

{ multifidelity_stoch_collocation
  [ { p_refinement
      uniform
      | ( dimension_adaptive
        sobol
      | generalized
    )
    ]
  ]
  [ { h_refinement
      uniform
      | ( dimension_adaptive
        sobol
      )
    ]
  ]
Keywords Area

| generalized |
| local_adaptive |

max_refinement_iterations INTEGER >= 0

convergence_tolerance REAL

metric_scale
relative
absolute

statistics_mode
active
combined

allocation_control
greedy

discrepancy_emulation
distinct ALIAS paired
recursive

quadrature_order_sequence INTEGERLIST

dimension_preference REALLIST |

nested
non_nested }

sparse_grid_level_sequence INTEGERLIST

dimension_preference REALLIST |

nodal
hierarchical |

restricted
unrestricted |

nested
non_nested }

piecewise
askey
wiener

use_derivatives |
samples_on_emulator ALIAS samples INTEGER |
sample_type
lhs
random |

rng
mt19937
rnum2 |

probability_refinement ALIAS sample_refinement import
adapt_import
mm_adapt_import

refinement_samples INTEGERLIST }

final_moments
none
standard
central |

response_levels REALLIST

num_response_levels INTEGERLIST |

compute
probabilities
reliabilities
gen_reliabilities
system
series
parallel }

probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
[
reliability_levels REALLIST
[ num_reliability_levels INTEGERLIST ]
[
gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
distribution
| cumulative
| complementary
[
variance_based_decomp
| interaction_order INTEGER > 0 |
| drop_tolerance REAL |
]
diagonal_covariance
| full_covariance |
import_approx_points_file STRING
[ ( custom.annotated
| header ]
| eval_id ]
[ interface_id ]
) |
annotated
| freeform |
| active_only |
[
export_approx_points_file ALIAS export_points_file STRING
[ ( custom.annotated
| header ]
| eval_id ]
[ interface_id ]
) |
annotated
| freeform |
| active_only |
]
seed_sequence INTEGERLIST
[ fixed_seed ]
[ model_pointer STRING ]
]
sampling ALIAS nond_sampling
[ samples ALIAS initial_samples INTEGER ]
[ seed INTEGER > 0 ]
[ fixed_seed ]
sample_type
| lhs |
| random |
| incremental_lhs |
| incremental_random |
[
refinement_samples INTEGERLIST ]
d_optimal
[ candidate_designs INTEGER > 0 |
| leja_oversample_ratio REAL ]
[
variance_based_decomp
| drop_tolerance REAL |
]
backfill
principal_components
[ percent.variance.explained REAL ]
wilks
[ order INTEGER ]
[ confidence_level REAL ]
[ one.sided.lower ]
[ one.sided.upper ]
| two.sided |
[ final.moments
| none ]
Keywords Area

response_levels REALLIST
  [ num_response_levels INTEGERLIST ]
  [ compute
    probabilities
    | reliabilities
    | gen_reliabilities
    | system
    series
    | parallel
  ]

probability_levels REALLIST
  [ num_probability_levels INTEGERLIST ]

reliability_levels REALLIST
  [ num_reliability_levels INTEGERLIST ]

gen_reliability_levels REALLIST
  [ num_gen_reliability_levels INTEGERLIST ]

distribution
  cumulative
  | complementary

rng
  mt19937
  | rnum2

model_pointer STRING

multilevel_sampling ALIAS multilevel_mc ALIAS mlmc
  [ seed_sequence INTEGERLIST ]
  [ fixed_seed ]
  [ pilot_samples ALIAS initial_samples INTEGERLIST ]
  [ sample_type
    lhs
    | random
  ]

export_sample_sequence
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  ]
  [ annotated
    | freeform
  ]

allocation_target
  mean
  |
  ( variance
    [ optimization ]
  )
  |
  ( standard_deviation
    [ optimization ]
  )
  |
  ( scalarization
    [ scalarization_response_mapping REALLIST ]
    [ optimization ]
  )

qoi_aggregation
  sum
  | max
  |
  [ convergence_tolerance REAL ]
[ convergence_tolerance_type
  relative
  | absolute
]
[ convergence_tolerance_target
  variance_constraint
  | cost_constraint
]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ final_moments
  none
  | standard
  | central
]
[ distribution
  cumulative
  | complementary
]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)

( multifidelity_sampling ALIAS multifidelity_mc ALIAS mfmc
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ pilot_samples ALIAS initial_samples INTEGERLIST ]
[ solution_mode
  online_pilot
  | offline_pilot
  | pilot_projection
]
[ sample_type
  lhs
  | random
]
[ export_sample_sequence
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
]
[ convergence_tolerance REAL ]
[ max_iterations INTEGER >= 0 ]
[ max_function_evaluations INTEGER >= 0 ]
[ final_moments
  none
  | standard
  | central
]
[ distribution
  cumulative
  | complementary
]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)

( multilevel_multifidelity_sampling ALIAS multilevel_multifidelity_mc ALIAS mlmfmc
[ seed_sequence INTEGERLIST ]
[ fixed_seed ]
[ pilot_samples ALIAS initial_samples INTEGERLIST ]
[ sample_type
  lhs
  | random
]
random

export_sample_sequence
  ( custom_annotated
    header
    eval_id
    interface_id
  )
  annotated
  freeform

convergence_tolerance REAL
max_iterations INTEGER >= 0
max_function_evaluations INTEGER >= 0

final_moments
  none
  standard
  central

distribution
  cumulative
  complementary

rng
  mt19937
  rnum2

model_pointer STRING

approximate_control_variate ALIAS acv_sampling
acv_independent_sampling ALIAS acv_is
acv_multifidelity ALIAS acv_mf
acv_kl ALIAS acv_adaptive
pilot_samples ALIAS initial_samples INTEGERLIST

solution_mode
  online_pilot
  offline_pilot
  pilot_projection

truth_fixed_by_pilot
sqp
nip

seed_sequence INTEGERLIST
fixed_seed

sample_type
  lhs
  random

evaluated
  ( custom_annotated
    header
    eval_id
    interface_id
  )
  annotated
  freeform

convergence_tolerance REAL
max_iterations INTEGER >= 0
max_function_evaluations INTEGER >= 0

final_moments
  none
  standard
  central

distribution
  cumulative
  complementary

rng
  mt19937
  rnum2
import importance_sampling ALIAS nond_importance_sampling
| samples ALIAS initial_samples INTEGER |
| seed INTEGER > 0 |
import adapt_import
| mm_adapt_import |
| refinement_samples INTEGERLIST |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| response_levels REALLIST |
| num_response_levels INTEGERLIST |
| compute probabilities |
| gen_reliabilities |
| system series |
| parallel |
|
| probability_levels REALLIST |
| num_probability_levels INTEGERLIST |
| gen_reliability_levels REALLIST |
| num_gen_reliability_levels INTEGERLIST |
| distribution cumulative |
| complementary |
| rng mt19937 |
rnum2 |
| model_pointer STRING |
|
| gpais ALIAS gaussian_process_adaptive_importance_sampling |
| build_samples ALIAS samples INTEGER |
| seed INTEGER > 0 |
| samples_on_emulator INTEGER |
| import_build_points_file ALIAS import_points_file STRING |
| ( custom_annotated |
| header ] |
| eval_id ] |
| interface_id ] |
| annotated |
| freeform |
| active_only ] |
|
| export_approx_points_file ALIAS export_points_file STRING |
| ( custom_annotated |
| header ] |
| eval_id ] |
| interface_id ] |
| annotated |
| freeform |
| ] |
| max_iterations INTEGER >= 0 |
| response_levels REALLIST |
| num_response_levels INTEGERLIST |
| compute probabilities |
| gen_reliabilities |
| system series |
| parallel |
}]
}
[probability_levels REALLIST
  [num_probability_levels INTEGERLIST]]
[gen_reliability_levels REALLIST
  [num_gen_reliability_levels INTEGERLIST]]
[distribution
  cumulative
  | complementary
  ]
[rng
  mt19937
  | rnum2
  ]
[model_pointer STRING]
)
)
(adaptive_sampling ALIAS nond_adaptive_sampling
[initial_samples ALIAS samples INTEGER]
[seed INTEGER > 0]
samples_on_emulator INTEGER]
[fitness_metric
  predicted_variance
  | distance
  | gradient
  ]
batch_selection
  naive
  | distance_penalty
  | topology
  | constant_liar
  ]
[refinement_samples INTEGERLIST]
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    [eval_id]
    [interface_id]
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  | freeform
  | active_only]
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    [eval_id]
    [interface_id]
  ]
  | annotated
  | freeform
  ]
)[misc_options STRINGLIST]
[max_iterations INTEGER >= 0]
[response_levels REALLIST
  [num_response_levels INTEGERLIST]
  [compute
    probabilities
    | gen_reliabilities
    | system
    series
    | parallel
    ]
  ]
)[probability_levels REALLIST
  [num_probability_levels INTEGERLIST]]
[gen_reliability_levels REALLIST
  [num_gen_reliability_levels INTEGERLIST]]
7.3 model

) [ distribution
cumulative
| complementary
] [ rng
mt19937
| rnum2
] [ model_pointer STRING ] )

[ pof_darts ALIAS nond_pof_darts
build_samples ALIAS samples INTEGER
[ seed INTEGER > 0 ]
[ lipschitz
local
| global
] [ samples_on_emulator INTEGER ] [ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| gen_reliabilities
| system
series
| parallel
]
] [ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
] [ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
] [ distribution
cumulative
| complementary
] [ rng
mt19937
| rnum2
] [ model_pointer STRING ] )

[ rkd_darts ALIAS nond_rkd_darts
build_samples ALIAS samples INTEGER
[ seed INTEGER > 0 ]
[ lipschitz
local
| global
] [ samples_on_emulator INTEGER ] [ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
| probabilities
| gen_reliabilities
| system
series
| parallel
]
] [ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
] [ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
cumulative
| complementary 
]
[ rng
mt19937
| rnum2
]
[ model_pointer STRING ]
)

( global_evidence ALIAS nond_global_evidence
[ samples INTEGER ]
[ seed INTEGER > 0 ]
( sbo
[ gaussian_process ALIAS kriging
surfpack
| dakota
]
[ use_derivatives ]
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform
| active_only
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform
]
)

( ego
gaussian_process ALIAS kriging
surfpack
| dakota
]
[ use_derivatives ]
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform
| active_only
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
) ]
| annotated
| freeform
]
)
| ea
| lhs ]
[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
probabilities
]
| gen_reliabilities
| system
| series
| parallel
|
|
| probability_levels REALIST
| num_probability_levels INTEGERLIST |
|
| gen_reliability_levels REALIST
| num_gen_reliability_levels INTEGERLIST |
|
| distribution
| cumulative
| complementary
|
| rng
| mt19937
| rnum2
|
| model_pointer STRING |
|
| global_interval_est ALIAS nond_global_interval_est
| samples INTEGER |
| seed INTEGER > 0 |
| max_iterations INTEGER >= 0 |
| convergence_tolerance REAL |
| max_function_evaluations INTEGER >= 0 |
| ( sbo
| |
| | gaussian_process ALIAS kriging
| | surfpack
| | dakota
|
| | ( experimental
| | | options_file STRING |
| | )
|
| | use_derivatives |
| | import_build_points_file ALIAS import_points_file STRING |
| | ( custom_annotated
| | | header |
| | | eval_id |
| | | interface_id |
| | )
| | annotated
| | freeform |
| | active_only |
| | export_approx_points_file ALIAS export_points_file STRING |
| | ( custom_annotated
| | | header |
| | | eval_id |
| | | interface_id |
| | )
| | annotated
| | freeform |
| |
|
| ( ego
| | gaussian_process ALIAS kriging
| | surfpack
| | export_model
| | | filename_prefix STRING |
| | | formats
| | | text_archive |
| | | binary_archive |
| | )
| } dakota
( experimental
[ export_model
  [ filename_prefix STRING ]
  [ formats
    [ text_archive ]
    [ binary_archive ]
  ]
]
[ options_file STRING ]
)

[ use_derivatives ]
[ import_build_points_file ALIAS import_points_file STRING
  [ { custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  }]
  | annotated
  | freeform
  | active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
  [ { custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  }]
  | annotated
  | freeform ]
]
[ ea
  | lhs ]
[ rng
  mt19937
  | rnum2
]
[ model_pointer STRING ]
)

( bayes_calibration ALIAS nond_bayes_calibration
  [ queso
    chain_samples ALIAS samples INTEGER
    [ seed INTEGER > 0 ]
    [ rng
      mt19937
      | rnum2
    ]
    [ emulator
      ( gaussian_process ALIAS kriging
        surfpack
        | dakota
        [ build_samples INTEGER ]
        [ posterior_adaptive ]
    ]
    [ import_build_points_file ALIAS import_points_file STRING
      [ { custom_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
      }]
      | annotated
      | freeform
      | active_only ]
  ]
)

( pce
  [ p_refinement
    uniform
    | ( dimension_adaptive

sobol
| decay
| generalized
|
|
[ max_refinement_iterations INTEGER >= 0 ]
{ quadrature_order INTEGER
  [ dimension_preference REALLIST ]
  [ nested
  | non_nested ]
}
|
{ sparse_grid_level INTEGER
  [ dimension_preference REALLIST ]
  [ restricted
  | unrestricted ]
  [ nested
  | non_nested ]
}
| cubature_integrand INTEGER
|
{ expansion_order INTEGER
  [ dimension_preference REALLIST ]
  [ basis_type
    tensor_product
    | total_order
    |
    adapted
    [ advancements INTEGER ]
    [ soft_convergence_limit INTEGER ]
  ]
}
|
{ collocation_points INTEGER
  [ ( least_squares
    | svd
    | equality_constrained ]
  )
  |
  [ orthogonal_matching_pursuit ALIAS omp
    [ noise_tolerance REALLIST ]
  ]
  [ basis_pursuit ALIAS bp
  ]
  [ basis_pursuit_denoising ALIAS bpdn
    [ noise_tolerance REALLIST ]
  ]
  ]
  |
  [ least_angle_regression ALIAS lars
    [ noise_tolerance REALLIST ]
  ]
  |
  [ least_absolute_shrinkage ALIAS lasso
    [ noise_tolerance REALLIST ]
    [ l2_penalty REAL ]
  ]
  [ cross_validation
    [ noise_only ]
    [ max_cv_order_candidates INTEGER >= 0 ]
  ]
  [ ratio_order REAL ]
  [ response_scaling ]
  [ use_derivatives ]
  [ tensor_grid ]
  [ reuse_points ALIAS reuse_samples ]
  [ max_solver_iterations INTEGER >= 0 ]
  )
|
{ collocation_ratio REAL
  [ ( least_squares
    | svd
    | equality_constrained ]
  )
  |
{ orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
}
| basis_pursuit ALIAS bp
| { basis_pursuit_denoising ALIAS bpdn
  [ noise_tolerance REALLIST ]
}
| { least_angle_regression ALIAS lars
  [ noise_tolerance REALLIST ]
}
| { least_absolute_shrinkage ALIAS lasso
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]
  [ cross_validation
    [ noise_only ]
    [ max_cv_order_candidates INTEGER >= 0 ]
  ]
  [ ratio_order REAL ]
  [ response_scaling ]
  [ use_derivatives ]
  [ tensor_grid ]
  [ reuse_points ALIAS reuse_samples ]
  [ max_solver_iterations INTEGER >= 0 ]
}
| { expansion_samples INTEGER
  [ reuse_points ALIAS reuse_samples ]
}
| [ import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ]
]
| [ posterior_adaptive ]
}
| { orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points INTEGER
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
  [ import_build_points_file ALIAS import_points_file STRING
    [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
  | annotated
  | freeform ]
  [ active_only ]
]
| [ posterior_adaptive ]
}
[ askey
| wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance ]
[ full_covariance ]
}
| { ml_pce
  allocation_control
  [ estimator_variance
    [ estimator_rate REAL ]
  ]
7.3 model

) [ rip_sampling ]
[ discrepancy_emulation
  distinct ALIAS paired
  recursive ]
{ expansion_order_sequence INTEGERLIST
  [ dimension_preference REALLIST ]
  [ basis_type
tensor_product
  | total_order
  |
  [ adapted
    [ advancements INTEGER ]
    [ soft_convergence_limit INTEGER ]
  ]
]
]
[ collocation_ratio REAL
  [ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
  [ ( least_squares
    [ svd
      | equality_constrained ]
  ]
  |
  { orthogonal_matching_pursuit ALIAS omp
    [ noise_tolerance REALLIST ]
  ]
  [ basis_pursuit ALIAS bp
  ]
  [ basis_pursuit_denoising ALIAS bpdn
    [ noise_tolerance REALLIST ]
  ]
  |
  { least_angle_regression ALIAS lars
    [ noise_tolerance REALLIST ]
  ]
  |
  { least_absolute_shrinkage ALIAS lasso
    [ noise_tolerance REALLIST ]
    [ l2_penalty REAL ]
  ]
  [ cross_validation
    [ noise_only ]
    [ max_cv_order_candidates INTEGER >= 0 ]
  ]
  [ ratio_order REAL ]
  [ response_scaling ]
  [ use_derivatives ]
  [ tensor_grid ]
  [ reuse_points ALIAS reuse_samples ]
  [ max_solver_iterations INTEGER >= 0 ]
]
]
( expansion_samples_sequence INTEGERLIST
  [ reuse_points ALIAS reuse_samples ]
)
[ import_build_points_file ALIAS import_points_file STRING
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    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ]
]
]
{ orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
collocation_points_sequence INTEGERLIST
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
}
[ import_build_points_file ALIAS import_points_file STRING
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    [ eval_id ]
    [ interface_id ]
  )]
  [ annotated
    [ freeform ]
  ]
  [ active_only ]
]
[ askey
  [ wiener ]
  [ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance
  | full_covariance ]
]
[ mf_pce
[ p_refinement
  uniform
  [
    ( dimension_adaptive
      sobol
      | decay
      | generalized
    )
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[ allocation_control
  greedy
]
[ discrepancy_emulation
  distinct ALIAS paired
  | recursive
]
[ quadrature_order_sequence INTEGERLIST
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  [ nested
    | non_nested ]
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    | unrestricted ]
  [ nested
    | non_nested ]
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    | total_order
    [
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      [ soft_convergence_limit INTEGER ]
    ]
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    [ svd
     | equality_constrained ]
  )]
[ orthogonal_matching_pursuit ALIAS omp
  [ noise_tolerance REALLIST ]
]
[ basis_pursuit ALIAS bp
]
7.3 model

| basis_pursuit_denoising ALIAS bpdn
  | noise_tolerance REALIST |
| least_angle_regression ALIAS lars
  | noise_tolerance REALIST |
| least_absolute_shrinkage ALIAS lasso
  | noise_tolerance REALIST |
  | l2_penalty REAL |
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    | noise_only |
    | max_cv_order_candidates INTEGER >= 0 |
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  | response_scaling |
  | use_derivatives |
  | tensor_grid |
  | reuse_points ALIAS reuse_samples |
  | max_solver_iterations INTEGER >= 0 |
  | response_scaling |
  | use_derivatives |
  | tensor_grid |
  | reuse_points ALIAS reuse_samples |
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    | noise_only |
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  | ratio_order REAL |
  | response_scaling |
  | use_derivatives |
  | tensor_grid |
  | reuse_points ALIAS reuse_samples |
    | noise_only |
    | max_cv_order_candidates INTEGER >= 0 |
| orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST |
  | reuse_points ALIAS reuse_samples |
  | import_build_points_file ALIAS import_points_file STRING |
  | ( custom_annotated
    | header |
    | eval_id |
    | interface_id |
    | annotated |
    | freeform |
    | active_only |
  |
| orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST |
  | reuse_points ALIAS reuse_samples |
  | import_build_points_file ALIAS import_points_file STRING |
  | ( custom_annotated
    | header |
    | eval_id |
    | interface_id |
    | annotated |
    | freeform |
    | active_only |
  |
| askey |
| wiener |
  | normalized |
  | export_expansion_file STRING |
  | diagonal_covariance |
  | full_covariance |
  | diagonal_covariance |
  | full_covariance |
| p_refinement
  | uniform |
  | ( dimension_adaptive
    | sobol |
    | generalized |
  |
| granted |
(h_refinement
uniform
|
(dimension_adaptive
  sobol
  | generalized
) | local_adaptive
}
[ max_refinement_iterations INTEGER >= 0 ]
)
quadrature_order INTEGER
[ dimension_preference REALLIST ]
[ nested | non_nested ]
)
|
(sparse_grid_level INTEGER
[ dimension_preference REALLIST ]
[ nodal | hierarchical ]
[ restricted | unrestricted ]
[ nested | non_nested ]
)
| piecewise
| askey
| wiener |
| use_derivatives |
| diagonal_covariance
| full_covariance |
)
|
(mf_sc
|
(p_refinement
uniform
|
(dimension_adaptive
  sobol
  | generalized
) |
)
|
(h_refinement
uniform
|
(dimension_adaptive
  sobol
  | generalized
) | local_adaptive
]
[ max_refinement_iterations INTEGER >= 0 ]
[ allocation_control
  greedy ]
[ discrepancy_emulation
  distinct ALIAS paired
  | recursive ]
)
quadrature_order_sequence INTEGERLIST
[ dimension_preference REALLIST ]
[ nested | non_nested ]
)
|
(sparse_grid_level_sequence INTEGERLIST
[ dimension_preference REALLIST ]
[ nodal | hierarchical ]
| restricted | unrestricted ]
| nested
)}
[ export_chain_points_file STRING
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  [ eval_id ]
  [ interface_id ]
}
| annotated
| freeform ]
]
dram
| delayed_rejection
| adaptive_metropolis
| metropolis_hastings ]
[ proposal_covariance
  ( prior
  [ multiplier REAL > 0.0 ]
)
| derivatives
| update_period INTEGER ]
)
| ( values REALLIST
  diagonal
  | matrix
|)
| ( filename STRING
  diagonal
  | matrix
|)
[ options_file STRING ]
)
|
wasabi
| pushforward_samples INTEGER
[ seed INTEGER > 0 ]
| emulator
  ( gaussian_process ALIAS kriging
  surfpack
  | dakota
  [ build_samples INTEGER ]
  [ posterior_adaptive ]
| import_build_points_file ALIAS import_points_file STRING
[ { custom_annotated
  [ header ]
  [ eval_id ]
  [ interface_id ]
}
| annotated
| freeform ]
| active_only ]
)
)
| pce
[ p_refinement
  uniform
  |
  ( dimension_adaptive
  | sobol
  | decay
  | generalized
  )
]
[ max_refinement_iterations INTEGER >= 0 ]
| quadrature_order INTEGER
  [ dimension_preference REALLIST ]
  | nested
  | non_nested ]
|)
( sparse_grid_level INTEGER
  [ dimension_preference REALLIST ]
  [ restricted ]
  [ unrestricted ]
  [ nested ]
  [ non_nested ]
 )
| cubature_integrand INTEGER |
| expansion_order INTEGER |
[ dimension_preference REALLIST ]
[ basis_type tensor_product |
  total_order |
  [ adapted |
    [ advancements INTEGER ]
    [ soft_convergence_limit INTEGER ]
  ] |
| collocation_points INTEGER |
  [ ( least_squares |
    [ svd |
    [ equality_constrained ]
  ) |
  [ orthogonal_matching_pursuit ALIAS omp |
    [ noise_tolerance REALLIST ]
  ]
| basis_pursuit ALIAS bp |
| basis_pursuit_denoising ALIAS bpdn |
  [ noise_tolerance REALLIST ]
)|
| least_angle_regression ALIAS lars |
  [ noise_tolerance REALLIST ]
)|
| least_absolute_shrinkage ALIAS lasso |
  [ noise_tolerance REALLIST ]
  [ l2_penalty REAL ]
)|
  [ cross_validation |
    [ noise_only ]
    [ max_cv_order_candidates INTEGER >= 0 ]
  ]
| ratio_order REAL |
| response_scaling |
| use_derivatives |
| tensor_grid |
| reuse_points ALIAS reuse_samples |
  [ max_solver_iterations INTEGER >= 0 ]
)|
| collocation_ratio REAL |
  [ ( least_squares |
    [ svd |
    [ equality_constrained ]
  ) |
  [ orthogonal_matching_pursuit ALIAS omp |
    [ noise_tolerance REALLIST ]
  ]
| basis_pursuit ALIAS bp |
| basis_pursuit_denoising ALIAS bpdn |
  [ noise_tolerance REALLIST ]
)|
| least_angle_regression ALIAS lars |
  [ noise_tolerance REALLIST ]
)
least_absolute_shrinkage ALIAS lasso
[ noise_tolerance REALIST ]
[ l2_penalty REAL ]
)
[ cross_validation
[ noise_only ]
[ max_cv_order_candidates INTEGER >= 0 ]
)
[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]
)
)
( expansion_samples INTEGER
[ reuse_points ALIAS reuse_samples ]
)
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
[ active_only ]
)
[ posterior_adaptive ]
)
)
( orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
[ collocation_points INTEGER
[ tensor_grid INTEGERLIST ]
[ reuse_points ALIAS reuse_samples ]
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
[ active_only ]
)
[ posterior_adaptive ]
)
)
[ askey
| wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance
| full_covariance ]
)
)
( ml_pce
[ allocation_control
[ estimator_variance
[ estimator_rate REAL ]
]
| rip_sampling
]
[ discrepancy_emulation
distinct ALIAS paired
| recursive
)
[ expansion_order_sequence INTEGERLIST
[ dimension_preference REALIST ]
[ basis_type
[ tensor_product
| total_order

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7.3 model

| adapted
| [ advancements INTEGER ]
| [ soft_convergence_limit INTEGER ]
|
| [ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
| [ ( least_squares
| [ svd
| [ equality_constrained ] ) ]
| { orthogonal_matching_pursuit ALIAS omp
| [ noise_tolerance REALLIST ] ]
| basis_pursuit ALIAS bp
| [ basis_pursuit_denoising ALIAS bpdn
| [ noise_tolerance REALLIST ] ]
| least_angle_regression ALIAS lars
| [ noise_tolerance REALLIST ]
| least_absolute_shrinkage ALIAS lasso
| [ noise_tolerance REALLIST ]
| [ l2_penalty REAL ]
| [ cross_validation
| [ noise_only ]
| [ max_cv_order_candidates INTEGER GEQ 0 ] ]
| [ ratio_order REAL ]
| [ response_scaling ]
| [ use_derivatives ]
| [ tensor_grid ]
| [ reuse_points ALIAS reuse_samples ]
| [ max_solver_iterations INTEGER GEQ 0 ]
|
| [ expansion_samples_sequence INTEGERLIST
| [ reuse_points ALIAS reuse_samples ] ]
|
| [ import_build_points_file ALIAS import_points_file STRING
| [ ( custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ] )
| annotated
| [ freeform ]
| [ active_only ] ]
|
| [ orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
| collocation_points_sequence INTEGERLIST ]
| [ tensor_grid INTEGERLIST ]
| [ reuse_points ALIAS reuse_samples ]
| [ import_build_points_file ALIAS import_points_file STRING
| [ ( custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ] )
| annotated
| [ freeform ]
| [ active_only ] ]
|
| [ askey

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wiener
normalized
export_expansion_file STRING
diagonal_covariance
full_covariance
]

dimension_adaptive
sobol
decay
generalized
)

max_refinement_iterations INTEGER >= 0

allocation_control
greedy

discrepancy_emulation
distinct ALIAS paired
recursive

quadrature_order_sequence INTEGERLIST
dimension_preference REALLIST
nested
non_nested

sparse_grid_level_sequence INTEGERLIST
dimension_preference REALLIST
restricted
unrestricted
nested
non_nested

expansion_order_sequence INTEGERLIST
dimension_preference REALLIST
basis_type
tensor_product
total_order
adapted
advancements INTEGER
soft_convergence_limit INTEGER

collection_ratio REAL
collection_points_sequence ALIAS pilot_samples INTEGERLIST
(least_squares
svd
equality_constrained
)

orthogonal_matching_pursuit ALIAS omp
noise_tolerance REALLIST

basis_pursuit ALIAS bp

basis_pursuit_denoising ALIAS bpdn
noise_tolerance REALLIST

least_angle_regression ALIAS lars
noise_tolerance REALLIST

least_absolute_shrinkage ALIAS lasso
noise_tolerance REALLIST
l2_penalty REAL

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[ cross_validation
  [ noise_only ]
  [ max_cv_order_candidates INTEGER >= 0 ]
]
[ ratio_order REAL ]
[ response_scaling ]
[ use_derivatives ]
[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]
)
|
( expansion_samples_sequence INTEGERLIST
  [ reuse_points ALIAS reuse_samples ]
)
[ import_build_points_file ALIAS import_points_file STRING
  [ ( custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  )
  | annotated
  | freeform ]
  [ active_only ]
]
)
|
( orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
  collocation_points_sequence INTEGERLIST
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
  [ import_build_points_file ALIAS import_points_file STRING
    [ ( custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
    | annotated
    | freeform ]
    [ active_only ]
  ]
)
[ askey
  | wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance
  | full_covariance ]
)
|
( sc
  [ ( p_refinement
    uniform
    |
    { dimension_adaptive
      sobol
      | generalized
    }
  )
  | h_refinement
    uniform
    |
    { dimension_adaptive
      sobol
      | generalized
    |
    local_adaptive
  ]
( max_refinement_iterations INTEGER >= 0 ]
( quadrature_order INTEGER
  [ dimension_preference REALLIST ]
)
[ nested
 | non_nested ]
)

[ sparse_grid_level INTEGER
 | dimension_preference REALLIST ]
 | nodal
 | hierarchical ]
 | restricted
 | unrestricted ]
 | nested
 | non_nested ]
)

[ piecewise
 | askey
 | wiener ]
 | use_derivatives ]
 | diagonal_covariance
 | full_covariance ]
)

[ mf_sc
 | p_refinement
 | uniform ]
 | dimension_adaptive
 | sobol
 | generalized ]
)

[ h_refinement
 | uniform ]
 | dimension_adaptive
 | sobol
 | generalized ]
 | local_adaptive ]

[ max_refinement_iterations INTEGER >= 0 ]
| allocation_control
 | greedy ]
| discrepancy_emulation
 | distinct ALIAS paired
 | recursive ]

| quadrature_order_sequence INTEGERLIST
 | dimension_preference REALLIST ]
 | nested
 | non_nested ]
)

| sparse_grid_level_sequence INTEGERLIST
 | dimension_preference REALLIST ]
 | nodal
 | hierarchical ]
 | restricted
 | unrestricted ]
 | nested
 | non_nested ]
)

[ piecewise
 | askey
 | wiener ]
 | use_derivatives ]
 | diagonal_covariance
 | full_covariance ]
)

[ standardized_space ]
| data_distribution
(gaussian
means REALLIST
(covariance REALLIST
diagonal
| matrix
)
| obs_data_filename STRING
)
| posterior_samples_import_filename STRING ]
generate_posterior_samples
| posterior_samples_export_filename STRING ]
evaluate_posterior_density
| posterior_density_export_filename STRING ]
)
|
dream
chain_samples ALIAS samples INTEGER
| seed INTEGER > 0 ]
| chains INTEGER >= 3 ]
| num_cr INTEGER >= 1 ]
| crossover_chain_pairs INTEGER >= 0 ]
| gr_threshold REAL > 0.0 ]
| jump_step INTEGER >= 0 ]
| emulator
( gaussian_process ALIAS kriging
surfpack
| dakota
| build_samples INTEGER ]
| posterior_adaptive ]
| import_build_points_file ALIAS import_points_file STRING
| { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
}
| annotated
| freeform ]
| active_only ]
)
|
pce
| p_refinement
uniform
|
( dimension_adaptive
sobol
| decay
| generalized
)
]
| max_refinement_iterations INTEGER >= 0 ]
| quadrature_order INTEGER
| dimension_preference REALLIST ]
| nested
| non_nested ]
|
| sparse_grid_level INTEGER
| dimension_preference REALLIST ]
| restricted
| unrestricted ]
| nested
| non_nested ]
|
sparse_grid_level INTEGER
| dimension_preference REALLIST ]
| restricted
| unrestricted ]
| nested
| non_nested ]
|
cubature_integrand INTEGER
|
| expansion_order INTEGER
| dimension_preference REALLIST ]
| basis_type
tensor_product
| total_order
|
| [ adapted
| [ advancements INTEGER ]
| [ soft_convergence_limit INTEGER ]
| ]
|
| [ collocation_points INTEGER
| [ [ least_squares
| [ svd
| [ equality_constrained ]
| ]
| [ orthogonal_matching_pursuit ALIAS omp
| [ noise_tolerance REALLIST ]
| ]
| [ basis_pursuit ALIAS bp
| [ basis_pursuit_denoising ALIAS bpdn
| [ noise_tolerance REALLIST ]
| ]
| [ least_angle_regression ALIAS lars
| [ noise_tolerance REALLIST ]
| ]
| [ least_absolute_shrinkage ALIAS lasso
| [ noise_tolerance REALLIST ]
| [ l2_penalty REAL ]
| ]
| [ cross_validation
| [ noise_only ]
| [ max_cv_order_candidates INTEGER >= 0 ]
| ]
| [ ratio_order REAL ]
| [ response_scaling ]
| [ use_derivatives ]
| [ tensor_grid ]
| [ reuse_points ALIAS reuse_samples ]
| [ max_solver_iterations INTEGER >= 0 ]
| ]
|
| [ collocation_ratio REAL
| [ [ least_squares
| [ svd
| [ equality_constrained ]
| ]
| [ orthogonal_matching_pursuit ALIAS omp
| [ noise_tolerance REALLIST ]
| ]
| [ basis_pursuit ALIAS bp
| [ basis_pursuit_denoising ALIAS bpdn
| [ noise_tolerance REALLIST ]
| ]
| [ least_angle_regression ALIAS lars
| [ noise_tolerance REALLIST ]
| ]
| [ least_absolute_shrinkage ALIAS lasso
| [ noise_tolerance REALLIST ]
| [ l2_penalty REAL ]
| ]
| [ cross_validation
| [ noise_only ]
| [ max_cv_order_candidates INTEGER >= 0 ]
| ]
| [ ratio_order REAL ]
| [ response_scaling ]
| [ use_derivatives ]

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[ tensor_grid ]
[ reuse_points ALIAS reuse_samples ]
[ max_solver_iterations INTEGER >= 0 ]
)
|
| ( expansion_samples INTEGER
[ reuse_points ALIAS reuse_samples ]
)

[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
[ active_only ]
]
[ posterior_adaptive ]
)
|
| ( orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
collocation_points INTEGER
[ tensor_grid INTEGERLIST ]
[ reuse_points ALIAS reuse_samples ]
[ import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
[ active_only ]
]
[ posterior_adaptive ]
)

[ askey
| wiener ]
[ normalized ]
[ export_expansion_file STRING ]
[ diagonal_covariance
| full_covariance ]
)
|
| ( ml_pce
[ allocation_control
( estimator_variance
[ estimator_rate REAL ]
)
| rip_sampling
]
| discrepancy_emulation
distinct ALIAS paired
| recursive
]
( expansion_order_sequence INTEGERLIST
[ dimension_preference REALLIST ]
[ basis_type
tensor_product
| total_order
]
( adapted
[ advancements INTEGER ]
[ soft_convergence_limit INTEGER ]
)
]

( collocation_ratio REAL
[ collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
[ ( least_squares
[ svd
| equality_constrained ]
)
| orthogonal_matching_pursuit ALIAS omp
| [ noise_tolerance REALLIST ]
|]
| basis_pursuit ALIAS bp
|]
| basis_pursuit_denoising ALIAS bpdn
| [ noise_tolerance REALLIST ]
|]
| least_angle_regression ALIAS lars
| [ noise_tolerance REALLIST ]
|]
| least_absolute_shrinkage ALIAS lasso
| [ noise_tolerance REALLIST ]
| [ l2_penalty REAL ]
|]
| cross_validation
| [ noise_only ]
| [ max_cv_order_candidates INTEGER >= 0 ]
|]
| [ ratio_order REAL ]
| [ response_scaling ]
| [ use_derivatives ]
| [ tensor_grid ]
| [ reuse_points ALIAS reuse_samples ]
| [ max_solver_iterations INTEGER >= 0 ]
|]
| expansion_samples_sequence INTEGERLIST
| [ reuse_points ALIAS reuse_samples ]
|]
| [ import_build_points_file ALIAS import_points_file STRING
| [ ( custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
| )
| annotated
| [ freeform ]
| [ active_only ]
| ]
|]
| orthogonal_least_interpolation ALIAS least_interpolation ALIAS oli
| collocation_points_sequence INTEGERLIST
| [ tensor_grid INTEGERLIST ]
| [ reuse_points ALIAS reuse_samples ]
| [ import_build_points_file ALIAS import_points_file STRING
| [ ( custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
| )
| annotated
| [ freeform ]
| [ active_only ]
| ]
|]
| askey
| [ wiener ]
| [ normalized ]
| [ export_expansion_file STRING ]
| [ diagonal_covariance ]
| [ full_covariance ]
|]
|]
| mf_pce
| [ p_refinement
| [ uniform
| [ ( dimension_adaptive
| [ ...
7.3 model

sobol
| decay
| generalized
|
|
[ max_refinement_iterations INTEGER >= 0 ]
| allocation_control
| greedy
|
| discrepancy_emulation
| distinct ALIAS paired
| recursive
|
{ quadrature_order_sequence INTEGERLIST
| dimension_preference REALIST ]
| nested
| non_nested ]
|
|
{ sparse_grid_level_sequence INTEGERLIST
| dimension_preference REALIST ]
| restricted
| unrestricted ]
| nested
| non_nested ]
|
|
{ expansion_order_sequence INTEGERLIST
| dimension_preference REALIST ]
| basis_type
| tensor_product
| total_order
|
| adapted
| advancements INTEGER ]
| soft_convergence_limit INTEGER ]
|
| collocation_ratio REAL
| collocation_points_sequence ALIAS pilot_samples INTEGERLIST ]
| ( least_squares
| svd
| equality_constrained ]
|
| ( orthogonal_matching_pursuit ALIAS omp
| noise_tolerance REALIST ]
|
| basis_pursuit ALIAS bp
|
| basis_pursuit_denoising ALIAS bpdn
| noise_tolerance REALIST ]
|
| least_angle_regression ALIAS lars
| noise_tolerance REALIST ]
|
| least_absolute_shrinkage ALIAS lasso
| noise_tolerance REALIST ]
| 12_penalty REAL ]
|
| cross_validation
| noise_only ]
| max_cv_order_candidates INTEGER >= 0 ]
|
| ratio_order REAL ]
| response_scaling ]
| use_derivatives ]
| tensor_grid ]
| reuse_points ALIAS reuse_samples ]
| max_solver_iterations INTEGER >= 0 ]
|

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( expansion_samples_sequence INTEGERLIST
  [ reuse_points ALIAS reuse_samples ]
)

[ import_build_points_file ALIAS import_points_file STRING
  [ { custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    } annotated
    [ freeform ]
    [ active_only ]
  ]
]

( orthogonal_least_interpolation ALIAS least_interpolation_ALIAS oli
  collocation_points_sequence INTEGERLIST
  [ tensor_grid INTEGERLIST ]
  [ reuse_points ALIAS reuse_samples ]
  [ import_build_points_file ALIAS import_points_file STRING
    [ { custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    } annotated
    [ freeform ]
    [ active_only ]
  ]
]

[ askey
  [ wiener ]
  [ normalized ]
  [ export_expansion_file STRING ]
  [ diagonal_covariance
    [ full_covariance ]
  ]]

( sc
  [ p_refinement
    uniform
    [
      { dimension_adaptive
        sobol
        [ generalized
        }
    ]
  ]

  [ h_refinement
    uniform
    [
      { dimension_adaptive
        sobol
        [ generalized
        ]
      [ local_adaptive
      ]
    ]
    [ max_refinement_iterations INTEGER >= 0 ]
    { quadrature_order INTEGER
      [ dimension_preference REALLIST ]
      [ nested
      [ non_nested ]
    ]
  ]

  [ sparse_grid_level INTEGER
    [ dimension_preference REALLIST ]
    [ nodal
    [ hierarchical ]
    [ restricted
    [ nested
    [ non_nested ]
  ]}
7.3 model

{ piecewise
| askey
| wiener }
{ use_derivatives }
{ diagonal_covariance
| full_covariance }

{ mf_sc
{ p_refinement
  uniform
  { dimension_adaptive
    sobol
    | generalized
  }
}
{ h_refinement
  uniform
  { dimension_adaptive
    sobol
    | generalized
  }
  local_adaptive
}
{ max_refinement_iterations INTEGER >= 0 }
{ allocation_control
  greedy }
{ discrepancy_emulation
  distinct ALIAS paired
  | recursive }
{ quadrature_order_sequence INTEGERLIST
  { dimension_preference REALLIST }
  nested
  | non_nested }

{ sparse_grid_level_sequence INTEGERLIST
  { dimension_preference REALLIST }
  nodal
  | hierarchical }
  restricted
  | unrestricted }
  nested
  | non_nested }
{ piecewise
| askey
| wiener }
{ use_derivatives }
{ diagonal_covariance
| full_covariance }
}
{ standardized_space }
{ export_chain_points_file STRING
  { custom_annotated
    header
  }
  eval_id
  interface_id
}
{ annotated
  | freeform }
}
{ muq
  chain_samples ALIAS samples INTEGER
[ seed INTEGER > 0 ]
[ rng
  mt19937
  | rnum2
 ]
[ dram
  | delayed_rejection
  | adaptive_metropolis
  | metropolis_hastings ]
[ proposal_covariance
  ( prior
    [ multiplier REAL > 0.0 ]
  )]
  ( derivatives
    [ update_period INTEGER ]
  )
  ( values REALLIST
    diagonal
    | matrix
  )
  ( filename STRING
    diagonal
    | matrix
  )
]
[ experimental_design
  initial_samples ALIAS samples INTEGER
  num_candidates INTEGER > 0
  [ max_hifi_evaluations INTEGER >= 0 ]
  [ batch_size INTEGER >= 1 ]
  [ import_candidate_points_file STRING
    { custom_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    } | annotated | freeform ]
  ]
  [ ksg2 ]
]
[ calibrate_error_multipliers
  one
  | per_experiment
  | per_response
  | both
  [ hyperprior_alphas REALLIST
    hyperprior_betas REALLIST
  ]
]
[ burn_in_samples INTEGER ]
[ posterior_stats
  [ kl_divergence ]
  [ mutual_info
    [ ksg2 ]
  ]
  [ kde ]
]
[ chain_diagnostics
  [ confidence_intervals ]
]
[ model_evidence
  [ mc_approx ]
  [ evidence_samples INTEGER ]
  [ laplace_approx ]
]
[ model_discrepancy
  [ discrepancy_type
    ( gaussian_process ALIAS kriging
    | ...
  )
  ]
]

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[ trend_order INTEGER >= 0 ]

| ( polynomial
| [ basis_order INTEGER >= 0 ]
|
| [ num_prediction_configs INTEGER >= 0 ]
| prediction_configs REALLIST ]
| import_prediction_configs STRING
| { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
|
| annotated
| freeform ]
|
| export_discrepancy_file STRING
| { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
|
| annotated
| freeform ]
|
| export_corrected_model_file STRING
| { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
|
| annotated
| freeform ]
|
| export_corrected_variance_file STRING
| { custom_annotated
| [ header ]
| [ eval_id ]
| [ interface_id ]
|
| annotated
| freeform ]
|
| sub_sampling_period INTEGER ]
| probability_levels REALLIST
| [ num_probability_levels INTEGERLIST ]
|
| convergence_tolerance REAL ]
| max_iterations INTEGER >= 0 ]
| model_pointer STRING ]
| [ scaling ]
|
| ( dace
| grid
| random
| oas
| lhs
| oa_lhs
| box_behnken
| central_composite
| [ samples INTEGER ]
| [ seed INTEGER > 0 ]
| [ fixed_seed ]
| [ main_effects ]
| [ quality_metrics ]
| [ variance_based_decomp
| [ drop_tolerance REAL ]
|
| symbols INTEGER ]
[ model_pointer STRING ]
)
]
[ fsu_cvt
[ samples INTEGER ]
[ seed INTEGER > 0 ]
[ fixed_seed ]
[ latinize ]
[ quality_metrics ]
[ variance_based_decomp
[ drop_tolerance REAL ]
]
[ trial_type
grid
| halton
| random
]
[ num_trials INTEGER ]
[ max_iterations INTEGER >= 0 ]
[ model_pointer STRING ]
)
]
[ psuade_moat
[ partitions INTEGERLIST ]
[ samples INTEGER ]
[ seed INTEGER > 0 ]
[ model_pointer STRING ]
)
]
[ local_evidence ALIAS nond_local_evidence
[ sgp
| nip ]
[ response_levels REALLIST
[ num_response_levels INTEGERLIST ]
[ compute
  probabilities
  | gen_reliabilities
  ]
[ system
  series
  | parallel
  ]
]
[ probability_levels REALLIST
[ num_probability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
cumulative
| complementary
]
[ model_pointer STRING ]
)
]
[ local_interval_est ALIAS nond_local_interval_est
[ sgp
| nip ]
[ convergence_tolerance REAL ]
[ model_pointer STRING ]
)
]
[ local_reliability ALIAS nond_local_reliability
[ mpp_search
  x_taylor_mean
  | u_taylor_mean
  | x_taylor_mpp
  | u_taylor_mpp
  | x_two_point
  | u_two_point
  | x_multi_point
  | u_multi_point
]
7.3 model

[ no_approx
  | sqp
  | nip ]
[ integration
  | first_order
  | second_order
  | probability_refinement ALIAS sample_refinement
  | import
  | adapt_import
  | mm_adapt_import
  | refinement_samples INTEGERLIST ]
[ seed INTEGER > 0 ]
]
]
]
]
[ response_levels REALLIST
  | num_response_levels INTEGERLIST ]
[ compute
  | probabilities
  | reliabilities
  | gen_reliabilities
  | system
  | series
  | parallel
  ]
]
]
[ probability_levels REALLIST
  | num_probability_levels INTEGERLIST ]
]
[ reliability_levels REALLIST
  | num_reliability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALLIST
  | num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
  | cumulative
  | complementary
  ]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ final_moments
  | none
  | standard
  | central
  ]
[ model_pointer STRING ]
]
]
[ global_reliability ALIAS nond_global_reliability
  | initial_samples INTEGER ]
[ x_gaussian_process ALIAS x_kriging
  | u_gaussian_process ALIAS u_kriging
  | surfpack
  | export_model
  | filename_prefix STRING ]
[ formats
  | text_archive
  | binary_archive ]
]
]
]
[ dakota
]
[ experimental
  | export_model
  | filename_prefix STRING ]
[ formats
  | text_archive
  | binary_archive ]
]
}
[ options_file STRING ]
[
import_build_points_file ALIAS import_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
| active_only ]
]
[ export_approx_points_file ALIAS export_points_file STRING
[ ( custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
)
| annotated
| freeform ]
]
[ use_derivatives ]
[ seed INTEGER > 0 ]
[rng
mt19937
| rnum2
]
[ response_levels REALIST
[ num_response_levels INTEGERLIST ]
| compute
probabilities
| gen_reliabilities
| system
series
| parallel
]
]
[ probability_levels REALIST
[ num_probability_levels INTEGERLIST ]
]
[ gen_reliability_levels REALIST
[ num_gen_reliability_levels INTEGERLIST ]
]
[ distribution
cumulative
| complementary
]
[ max_iterations INTEGER >= 0 ]
[ convergence_tolerance REAL ]
[ model_pointer STRING ]
]
[ fsu_quasi_mc
halton
| hammersley
| latinize
[ quality_metrics
| variance_based_decomp
[ drop_tolerance REAL ]
]
[ samples INTEGER ]
| fixed_sequence ]
[ sequence_start INTEGERLIST ]
| sequence_leap INTEGERLIST ]
| prime_base INTEGERLIST ]
| max_iterations INTEGER >= 0 ]
[ model_pointer STRING ]
]
[ vector_parameter_study
final_point REALIST
| step_vector REALIST
]
num_steps INTEGER
  [ model_pointer STRING ]
)

| ( list_parameter_study
  list_of_points REALIST
  |
  ( import_points_file STRING
    [ ( custom_ann_annotated
      [ header ]
      [ eval_id ]
      [ interface_id ]
    )
    | annotated
    | freeform
    [ active_only ]
  )
  [ model_pointer STRING ]
)

| ( centered_parameter_study
  step_vector REALIST
  steps_per_variable ALIAS deltas_per_variable INTEGERLIST
  [ model_pointer STRING ]
)

| ( multidim_parameter_study
  partitions INTEGERLIST
  [ model_pointer STRING ]
)

| ( richardson_extrap
  estimate_order
  | converge_order
  | converge_qoi
  [ refinement_rate REAL ]
  [ convergence_tolerance REAL ]
  [ max_iterations INTEGER >= 0 ]
  [ model_pointer STRING ]
)

KEYWORD model
  [ id_model STRING ]
  ( single ALIAS simulation
    [ interface_pointer STRING ]
    [ solution_level_cost REALIST
      [ solution_level_control STRING ]
    ]
  )

| ( surrogate
  [ id_surrogates INTEGERLIST ]
  [ global
    ( experimental_gaussian_process
      [ trend
        none
      | constant
      | linear
      | reduced_quadratic
      | quadratic
    )
    [ num_restarts INTEGER > 1 ]
    [ nugget REAL > 0 ]
    [ find_nugget INTEGER ]
    [ options_file STRING ]
    [ export_approx_variance_file STRING
      [ ( custom_ann_annotated
        [ header ]
        [ eval_id ]
        [ interface_id ]
      )
      | annotated
      | freeform
    )
  )

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[ export_model
[ filename_prefix STRING ]
( formats
[ text_archive ]
[ binary_archive ]
)
]
[ import_model
[ filename_prefix STRING ]
text_archive
[ binary_archive ]
]
)
( gaussian_process ALIAS kriging
( dakota
[ point_selection ]
[ trend
constant
| linear
| reduced_quadratic
]
)
)
( surfpack
[ trend
constant
| linear
| reduced_quadratic
| quadratic
]
)
[ optimization_method STRING ]
[ max_trials INTEGER > 0 ]
[ nugget REAL > 0 ]
[ find_nugget INTEGER ]
[ correlation_lengths REALLIST ]
[ export_model
[ filename_prefix STRING ]
( formats
[ text_archive ]
[ binary_archive ]
[ algebraic_file ]
[ algebraic_console ]
)
]
[ import_model
[ filename_prefix STRING ]
text_archive
[ binary_archive ]
]
[ export_approx_variance_file STRING
[ custom_annotated
[ header ]
[ eval_id ]
[ interface_id ]
]
[ annotated
[ freeform ]
]
)
)
( mars
[ max_bases INTEGER ]
[ interpolation
linear
| cubic
]
[ export_model
[ filename_prefix STRING ]
( formats
[ text_archive ]
[ binary_archive ]
)[ text_archive ]
[ binary_archive ]
)
[ binary_archive ]
)
)
[ import_model
[ filename_prefix STRING ]
text_archive
| binary_archive
] )
)
( moving_least_squares
[ basis_order ALIAS poly_order INTEGER >= 0 ]
[ weight_function INTEGER ]
[ export_model
[ filename_prefix STRING ]
(formats
| text_archive ]
| binary_archive ]
)
)[ import_model
[ filename_prefix STRING ]
text_archive
| binary_archive
] )
)
( function_train
| regression_type
ls
| ( rls2
l2_penalty REAL
)
)
[ max Solver_iterations INTEGER >= 0 ]
[ max Cross_iterations INTEGER >= 0 ]
[ solver_tolerance REAL ]
[ response_scaling ]
[ tensor_grid ]
[ rounding_tolerance REAL ]
[ arithmetic_tolerance REAL ]
[ start_order ALIAS order INTEGER >= 0
[ dimension_preference REALLIST ]
]
[ adapt_order ]
[ kick_order INTEGER > 0 ]
[ max_order INTEGER >= 0 ]
[ max_cv_order_candidates INTEGER >= 0 ]
[ start_rank ALIAS rank INTEGER >= 0 ]
[ adapt_rank ]
[ kick_rank INTEGER > 0 ]
[ max_rank INTEGER >= 0 ]
[ max_cv_rank_candidates INTEGER >= 0 ]
) )
)
( neural_network
[ max_nodes ALIAS nodes INTEGER ]
| range REAL ]
[ random_weight INTEGER ]
[ export_model
[ filename_prefix STRING ]
(formats
| text_archive ]
| binary_archive ]
| algebraic_file ]
| algebraic_console ]
)
)[ import_model
[ filename_prefix STRING ]
text_archive
| binary_archive
radial_basis

export_model

format

import_model

polynomial

export_model

import_model

experimental_polynomial

export_model

import_model

domain_decomposition

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold

jump_threshold
[ auto_refinement
  [ max_iterations INTEGER > 0 ]
  [ max_function_evaluations INTEGER > 0 ]
  [ convergence_tolerance REAL ]
  [ soft_convergence_limit INTEGER >= 0 ]
  [ cross_validation_metric STRING
    [ folds INTEGER > 0 ]
  ]
]

[ actual_model_pointer ALIAS truth_model_pointer STRING ]
[ reuse_points ALIAS reuse_samples
  all
  | region
  | none
]

[ import_build_points_file ALIAS import_points_file ALIAS samples_file STRING
  [ custom_annotated
    [ header
      [ use_variable_labels ]
    ]
    [ eval_id ]
    [ interface_id ]
  ]
  | ( annotated
    [ use_variable_labels ]
  ]
  | freeform ]
  [ active_only ]
]

[ export_approx_points_file ALIAS export_points_file STRING
  [ custom_annotated
    [ header ]
    [ eval_id ]
    [ interface_id ]
  ]
  | annotated
  | freeform ]
]

[ use_derivatives ]
[ correction
  zeroth_order
  | first_order
  | second_order
  additive
  | multiplicative
  | combined
]

[ metrics ALIAS diagnostics STRINGLIST
  [ cross_validation
    [ folds INTEGER
      | percent REAL ]
  ]
  [ press ]
]

[ import_challenge_points_file ALIAS challenge_points_file STRING
  [ custom_annotated
    [ header
      [ use_variable_labels ]
    ]
    [ eval_id ]
    [ interface_id ]
  ]
  | ( annotated
    [ use_variable_labels ]
  ]
  | freeform ]
  [ active_only ]
]
(multipoint
| qmea
  actual_model_pointer ALIAS truth_model_pointer STRING
  )

( local
taylor_series
  actual_model_pointer ALIAS truth_model_pointer STRING
  )

(hierarchical
  ordered_model_fidelities ALIAS model_fidelity_sequence STRINGLIST
    correction
      zeroth_order
      first_order
      second_order
      additive
      multiplicative
      combined
    )

(non_hierarchical ALIAS model_ensemble
  truth_model_pointer ALIAS actual_model_pointer STRING
  unordered_model_fidelities ALIAS approximation_models STRINGLIST
  )

(nested
  [ optional_interface_pointer STRING
    [ optional_interface_responses_pointer STRING ]
  ]

  [ sub_method_pointer STRING
    [ iterator_servers INTEGER > 0 ]
    [ iterator_scheduling
      master
      [ peer ]
    ]
    [ processors_per_iterator INTEGER > 0 ]
    [ primary_variable_mapping STRINGLIST ]
    [ secondary_variable_mapping STRINGLIST ]
    [ primary_response_mapping REALLIST ]
    [ secondary_response_mapping REALLIST ]
    [ identity_response_mapping ]
  ]

  [ active_subspace ALIAS subspace
    actual_model_pointer STRING
    [ initial_samples INTEGER ]
    [ sample_type
      lhs
      random
    ]
    [ truncation_method
      [ bing_li ]
      [ constantine ]
      [ energy
        [ truncation_tolerance REAL ]
      ]
      [ cross_validation
        [ minimum
          [ relative
            [ relative_tolerance REAL ]
            [ decrease
              [ relative_tolerance REAL ]
              [ decrease_tolerance REAL ]
              [ max_rank INTEGER ]
              [ exhaustive ]
            ]
          ]
        ]
      ]
      [ dimension INTEGER ]
      [ bootstrap_samples INTEGER ]
    ]
  )
7.3 model

\[ \text{build_surrogate} \]
\[ \text{refinement_samples INTEGERLIST } \]
\[ \text{normalization} \]
\[ \text{mean_value} \]
\[ \text{mean_gradient} \]
\[ \text{local_gradient} \]
\)
\(\text{adapted_basis} \)
\[ \text{actual_model_pointer STRING} \]
\[ \text{sparse_grid_level INTEGER} \]
\[ \text{expansion_order INTEGER} \]
\[ \text{collocation_ratio REAL} \]
\[ \text{dimension INTEGER} \]
\[ \text{rotation_method} \]
\[ \text{unranked} \]
\[ \text{ranked} \]
\[ \text{truncation_tolerance REAL} \]
\)
\(\text{random_field} \)
\[ \text{build_source} \]
\[ \text{rf_data_file STRING} \]
\[ \text{dace_method_pointer STRING} \]
\[ \text{analytic_covariance} \]
\[ \text{squared_exponential} \]
\[ \text{exponential} \]
\[ \text{expansion_form} \]
\[ \text{karhunen_loeve} \]
\[ \text{principal_components} \]
\[ \text{expansion_bases INTEGER} \]
\[ \text{truncation_tolerance REAL} \]
\[ \text{propagation_model_pointer STRING} \]
\[ \text{variables_pointer STRING} \]
\[ \text{responses_pointer STRING} \]
\[ \text{hierarchical_tagging} \]

\text{KEYWORD variables} \]
\[ \text{id_variables STRING } \]
\[ \text{active} \]
\[ \text{all} \]
\[ \text{design} \]
\[ \text{uncertain} \]
\[ \text{aleatory} \]
\[ \text{epistemic} \]
\[ \text{state} \]
\[ \text{mixed} \]
\[ \text{relaxed} \]
\[ \text{continuous_design INTEGER > 0} \]
\[ \text{initial_point ALIAS cdv_initial_point REALLIST } \]
\[ \text{lower_bounds ALIAS cdv_lower_bounds REALLIST } \]
\[ \text{upper_bounds ALIAS cdv_upper_bounds REALLIST } \]
\[ \text{scale_types ALIAS cdv_scale_types STRINGLIST } \]
\[ \text{scales ALIAS cdv_scales REALLIST } \]
\[ \text{descriptors ALIAS cdv_descriptors STRINGLIST } \]
\]
\[ \text{discrete_design_range INTEGER > 0} \]
\[ \text{initial_point ALIAS ddv_initial_point INTEGERLIST } \]
\[ \text{lower_bounds ALIAS ddv_lower_bounds INTEGERLIST } \]
\[ \text{upper_bounds ALIAS ddv_upper_bounds INTEGERLIST } \]
\[ \text{descriptors ALIAS ddv_descriptors STRINGLIST } \]
[ discrete_design_set
  [ integer INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values INTEGERLIST
    [ categorical STRINGLIST
      [ adjacency_matrix INTEGERLIST ]
    ]
    [ initial_point INTEGERLIST ]
    [ descriptors STRINGLIST ]
  ]
  [ string INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values STRINGLIST
    [ adjacency_matrix INTEGERLIST ]
    [ initial_point STRINGLIST ]
    [ descriptors STRINGLIST ]
  ]
  [ real INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values REALLIST
    [ categorical STRINGLIST
      [ adjacency_matrix INTEGERLIST ]
    ]
    [ initial_point REALLIST ]
    [ descriptors STRINGLIST ]
  ]
  [ normal_uncertain INTEGER > 0
    means ALIAS nuv_means REALLIST
    std_deviations ALIAS nuv_std_deviations REALLIST
    [ lower_bounds ALIAS nuv_lower_bounds REALLIST ]
    [ upper_bounds ALIAS nuv_upper_bounds REALLIST ]
    [ initial_point REALLIST ]
    [ descriptors ALIAS nuv_descriptors STRINGLIST ]
  ]
  [ lognormal_uncertain INTEGER > 0
    lambdas ALIAS lnuv_lambdas REALLIST
    zetas ALIAS lnuv_zetas REALLIST
    [ means ALIAS lnuv_means REALLIST
      std_deviations ALIAS lnuv_std_deviations REALLIST
      error_factors ALIAS lnuv_error_factors REALLIST ]
    [ lower_bounds ALIAS lnuv_lower_bounds REALLIST ]
    [ upper_bounds ALIAS lnuv_upper_bounds REALLIST ]
    [ initial_point REALLIST ]
    [ descriptors ALIAS lnuv_descriptors STRINGLIST ]
  ]
  [ uniform_uncertain INTEGER > 0
    lower_bounds ALIAS uuv_lower_bounds REALLIST
    upper_bounds ALIAS uuv_upper_bounds REALLIST
    [ initial_point REALLIST ]
    [ descriptors ALIAS uuv_descriptors STRINGLIST ]
  ]
  [ loguniform_uncertain INTEGER > 0
    lower_bounds ALIAS luuv_lower_bounds REALLIST
    upper_bounds ALIAS luuv_upper_bounds REALLIST
    [ initial_point REALLIST ]
    [ descriptors ALIAS luuv_descriptors STRINGLIST ]
  ]
  [ triangular_uncertain INTEGER > 0
    modes ALIAS tuv_modes REALLIST
    lower_bounds ALIAS tuv_lower_bounds REALLIST
    upper_bounds ALIAS tuv_upper_bounds REALLIST
    [ initial_point REALLIST ]
    [ descriptors ALIAS tuv_descriptors STRINGLIST ]
  ]
  [ exponential_uncertain INTEGER > 0
    betas ALIAS euv_betas REALLIST
    [ initial_point REALLIST ]
    [ descriptors ALIAS euv_descriptors STRINGLIST ]
]
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beta_uncertain INTEGER > 0
alphas ALIAS buv_alphas REALLIST
betas ALIAS buv_betas REALLIST
lower_bounds ALIAS buv_lower_bounds REALLIST
upper_bounds ALIAS buv_upper_bounds REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS buv_descriptors STRINGLIST ]

gamma_uncertain INTEGER > 0
alphas ALIAS gauv_alphas REALLIST
betas ALIAS gauv_betas REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS gauv_descriptors STRINGLIST ]

frechet_uncertain INTEGER > 0
alphas ALIAS fuv_alphas REALLIST
betas ALIAS fuv_betas REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS fuv_descriptors STRINGLIST ]

weibull_uncertain INTEGER > 0
alphas ALIAS wuv_alphas REALLIST
betas ALIAS wuv_betas REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS wuv_descriptors STRINGLIST ]

histogram_bin_uncertain INTEGER > 0
[ pairs_per_variable ALIAS num_pairs INTEGERLIST ]
abscissas ALIAS huv_bin_abscissas REALLIST
ordinates ALIAS huv_bin_ordinates REALLIST
counts ALIAS huv_bin_counts REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS huv_bin_descriptors STRINGLIST ]

poisson_uncertain INTEGER > 0
lambdas REALLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]

binomial_uncertain INTEGER > 0
probability_per_trial ALIAS prob_per_trial REALLIST
num_trials INTEGERLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]

negative_binomial_uncertain INTEGER > 0
probability_per_trial ALIAS prob_per_trial REALLIST
num_trials INTEGERLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]

geometric_uncertain INTEGER > 0
probability_per_trial ALIAS prob_per_trial REALLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]

hypergeometric_uncertain INTEGER > 0
total_population INTEGERLIST
selected_population INTEGERLIST
num_drawn INTEGERLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]

histogram_point_uncertain
[ integer INTEGER > 0
[ pairs_per_variable ALIAS num_pairs INTEGERLIST ]
abscissas INTEGERLIST
counts REALLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]
]
[ string INTEGER > 0
[ pairs_per_variable ALIAS num_pairs INTEGERLIST ]
abscissas STRINGLIST
counts REALLIST
[ initial_point STRINGLIST ]
[ descriptors STRINGLIST ]
]
[ real INTEGER > 0
[ pairs_per_variable ALIAS num_pairs INTEGERLIST ]
abscissas REALLIST
counts REALLIST
[ initial_point REALLIST ]
[ descriptors STRINGLIST ]
]
]
[ uncertain_correlation_matrix REALLIST ]
[ continuous_interval_uncertain ALIAS interval_uncertain INTEGER > 0
[ num_intervals ALIAS iuv_num_intervals INTEGERLIST ]
[ interval_probabilities ALIAS interval_probs ALIAS iuv_interval_probs REALLIST ]
upper_bounds REALLIST
[ initial_point REALLIST ]
[ descriptors ALIAS iuv_descriptors STRINGLIST ]
]
[ discrete_interval_uncertain INTEGER > 0
[ num_intervals INTEGERLIST ]
[ interval_probabilities ALIAS interval_probs ALIAS range_probabilities ALIAS range_probs REALLIST ]
lower_bounds INTEGERLIST
upper_bounds INTEGERLIST
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]
]
[ discrete_uncertain_set
[ integer INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values INTEGERLIST
[ set_probabilities ALIAS set_probs REALLIST ]
[ categorical STRINGLIST ]
[ initial_point INTEGERLIST ]
[ descriptors STRINGLIST ]
]
[ string INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values STRINGLIST
[ set_probabilities ALIAS set_probs REALLIST ]
[ categorical STRINGLIST ]
[ initial_point STRINGLIST ]
]
[ real INTEGER > 0
[ elements_per_variable ALIAS num_set_values INTEGERLIST ]
elements ALIAS set_values REALLIST
[ set_probabilities ALIAS set_probs REALLIST ]
[ categorical STRINGLIST ]
[ initial_point REALLIST ]
[ descriptors STRINGLIST ]
]
]
[ continuous_state INTEGER > 0
[ initial_state ALIAS csv_initial_state REALLIST ]
lower_bounds ALIAS csv_lower_bounds REALLIST
[ upper_bounds ALIAS csv_upper_bounds REALLIST ]
[ descriptors ALIAS csv_descriptors STRINGLIST ]
]
[ discrete_state_range INTEGER > 0
[ initial_state ALIAS dsv_initial_state INTEGERLIST ]
lower_bounds ALIAS dsv_lower_bounds INTEGERLIST
[ upper_bounds ALIAS dsv_upper_bounds INTEGERLIST ]
[ descriptors ALIAS dsv_descriptors STRINGLIST ]
]
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[ discrete_state_set
  [ integer INTEGER > 0
    [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
    elements ALIAS set_values INTEGERLIST
    categorical STRINGLIST ]
  [ initial_state INTEGERLIST ]
  [ descriptors STRINGLIST ]
]

[ string INTEGER > 0
  [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
  elements ALIAS set_values STRINGLIST
  initial_state STRINGLIST ]
  [ descriptors STRINGLIST ]
]

[ real INTEGER > 0
  [ elements_per_variable ALIAS num_set_values INTEGERLIST ]
  elements ALIAS set_values REALLIST
  initial_state REALLIST ]
  [ descriptors STRINGLIST ]
]

[ linear_inequality_constraint_matrix REALLIST ]
[ linear_inequality_lower_bounds REALLIST ]
[ linear_inequality_upper_bounds REALLIST ]
[ linear_inequality_scale_types STRINGLIST ]
[ linear_inequality_scales REALLIST ]
[ linear_equality_constraint_matrix REALLIST ]
[ linear_equality_targets REALLIST ]
[ linear_equality_scale_types STRINGLIST ]
[ linear_equality_scales REALLIST ]

KEYWORD interface
  [ id_interface STRING ]
  [ analysis_drivers STRINGLIST ]
  [ input_filter STRING ]
  [ output_filter STRING ]
  [ system
    [ parameters_file STRING ]
    [ results_file STRING ]
    [ file_tag ]
    [ file_save ]
    [ labeled ]
    [ aprepro ALIAS dprepro ]
  ]
  [ work_directory
    [ named STRING ]
    [ directory_tag ALIAS dir_tag ]
    [ directory_save ALIAS dir_save ]
    [ link_files STRINGLIST ]
    [ copy_files STRINGLIST ]
    [ replace ]
  ]
  [ allow_existing_results ]
  [ verbatim ]
]

[ fork
  [ parameters_file STRING ]
  [ results_file STRING ]
  [ file_tag ]
  [ file_save ]
  [ labeled ]
  [ aprepro ALIAS dprepro ]
  [ work_directory
    [ named STRING ]
    [ directory_tag ALIAS dir_tag ]
    [ directory_save ALIAS dir_save ]
    [ link_files STRINGLIST ]
    [ copy_files STRINGLIST ]
    [ replace ]
  ]
  [ allow_existing_results ]
]
Keywords Area

[ verbatim ]

| [ direct
  | processors_per_analysis INTEGER > 0 ]
  | matlab
  | pybind11
  | python
  | [ numpy ]
  | scilab
  | grid
  | [ analysis_components STRINGLIST ]
  |
  [ algebraic_mappings STRING ]

[ failure_capture
  abort
  | retry INTEGER
  | recover REALLIST
  | continuation
  |
  [ deactivate
    [ active_set_vector ]
    [ evaluation_cache ]
    [ strict_cache_equality
      [ cache_tolerance REAL ]
    ]
    [ restart_file ]
  ]

  | ( batch
    [ size INTEGER > 0 ]
  )

  | ( asynchronous
    [ evaluation_concurrency INTEGER > 0 ]
    [ local_evaluation_scheduling
      dynamic
      | static
    ]
    [ analysis_concurrency INTEGER > 0 ]
  ]

  | [ evaluation_servers INTEGER > 0 ]

[ evaluation_scheduling
  master
  | peer
    dynamic
    | static
  ]

| [ processors_per_evaluation INTEGER > 0 ]

[ analysis_servers INTEGER > 0 ]

[ analysis_scheduling
  master
  | peer

KEYWORD responses

[ id_responses STRING ]

[ descriptors ALIAS response_descriptors STRINGLIST ]

[ objective_functions ALIAS num_objective_functions INTEGER >= 0
  [ sense STRINGLIST ]
  [ primary_scale_types ALIAS objective_function_scale_types STRINGLIST ]
  [ primary_scales ALIAS objective_function_scales REALLIST ]
  [ weights ALIAS multi_objective_weights REALLIST ]
  [ nonlinear_inequality_constraints ALIAS num_nonlinear_inequality_constraints INTEGER >= 0
    [ lower_bounds ALIAS nonlinear_inequality_lower_bounds REALLIST ]
    [ upper_bounds ALIAS nonlinear_inequality_upper_bounds REALLIST ]
    [ scale_types ALIAS nonlinear_inequality_scale_types STRINGLIST ]
    [ scales ALIAS nonlinear_inequality_scales REALLIST ]
  ]

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[ nonlinear_equality_constraints ALIAS num_nonlinear_equality_constraints INTEGER >= 0 ]
[ targets ALIAS nonlinear_equality_targets STRINGLIST ]
[ scale_types ALIAS nonlinear_equality_scale_types STRINGLIST ]
[ scales ALIAS nonlinear_equality_scales REALLIST ]
]
[ scalar_objectives ALIAS num_scalar_objectives INTEGER >= 0 ]
[ field_objectives ALIAS num_field_objectives INTEGER >= 0 ]
lengths INTEGERLIST
[ num_coordinates_per_field INTEGERLIST ]
[ read_field_coordinates ]
]
)
]
[ scalar_calibration_terms INTEGER >= 0 ]
[ field_calibration_terms INTEGER >= 0 ]
lengths INTEGERLIST
[ num_coordinates_per_field INTEGERLIST ]
[ read_field_coordinates ]
]
[ primary_scales ALIAS calibration_term_scales ALIAS least_squares_term_scales REALLIST ]
[ weights ALIAS calibration_weights ALIAS least_squares_weights REALLIST ]
]
[ calibration_data ]
[ num_experiments INTEGER >= 0 ]
[ num_config_variables INTEGER >= 0 ]
[ experiment_variance_type ALIAS variance_type STRINGLIST ]
[ scalar_data_file STRING ]
( custom_annotated
[ header ]
[ exp_id ]
) | annotated
| freeform
)
[ interpolate ]
]
[ calibration_data_file ALIAS least_squares_data_file STRING ]
[ custom_annotated
[ header ]
[ exp_id ]
) | annotated
| freeform
[ num_experiments INTEGER >= 0 ]
[ num_config_variables INTEGER >= 0 ]
[ experiment_variance_type ALIAS variance_type STRINGLIST ]
] | simulation_variance REALLIST ]
[ nonlinear_inequality_constraints ALIAS num_nonlinear_inequality_constraints INTEGER >= 0 ]
[ lower_bounds ALIAS nonlinear_inequality_lower_bounds REALLIST ]
[ upper_bounds ALIAS nonlinear_inequality_upper_bounds REALLIST ]
[ scale_types ALIAS nonlinear_inequality_scale_types STRINGLIST ]
[ scales ALIAS nonlinear_inequality_scales REALLIST ]
]
[ nonlinear_inequality_constraints ALIAS num_nonlinear_inequality_constraints INTEGER >= 0 ]
[ targets ALIAS nonlinear_inequality_targets REALLIST ]
[ scale_types ALIAS nonlinear_inequality_scale_types STRINGLIST ]
[ scales ALIAS nonlinear_inequality_scales REALLIST ]
]
)
]
[ response_functions ALIAS num_response_functions INTEGER >= 0 ]
[ scalar_responses ALIAS num_scalar_responses INTEGER >= 0 ]
[ field_responses ALIAS num_field_responses INTEGER >= 0 ]
lengths INTEGERLIST
[ num_coordinates_per_field INTEGERLIST ]
[ read_field_coordinates ]
]
no_gradients
| analytic_gradients
|
(mixed_gradients
  id_numerical_gradients INTEGERLIST
  id_analytic_gradients INTEGERLIST
  [ method_source ]
  [ ( dakota
    [ ignore_bounds ]
    [ relative
      [ absolute
        [ bounds ]
      ]
    ]
    [ vendor ]
    [ interval_type ]
    [ forward
      [ central ]
    ]
    [ fd_step_size ALIAS fd_gradient_step_size REALLIST ]
  )]
  [ numerical_gradients
    [ method_source ]
    [ ( dakota
      [ ignore_bounds ]
      [ relative
        [ absolute
          [ bounds ]
        ]
      ]
      [ vendor ]
      [ interval_type ]
      [ forward
        [ central ]
      ]
      [ fd_step_size ALIAS fd_gradient_step_size REALLIST ]
    )]
  ]
  no_hessians
  [ numerical_hessians
    [ fd_step_size ALIAS fd_hessian_step_size REALLIST ]
    [ relative
      [ absolute
        [ bounds ]
      ]
    ]
    [ forward
      [ central ]
    ]
  ]
  [ quasi_hessians
    [ bfgs
      [ damped ]
    ]
    [ srl ]
  ]
  analytic_hessians
  [ mixed_hessians
    [ id_numerical_hessians INTEGERLIST
      [ fd_step_size ALIAS fd_hessian_step_size REALLIST ]
    ]
    [ relative
      [ absolute
        [ bounds ]
      ]
    ]
    [ forward
      [ central ]
    ]
    [ id_quasi_hessians INTEGERLIST
      [ bfgs
        [ damped ]
      ]
      [ srl ]
    ]
    [ id_analytic_hessians INTEGERLIST ]
  )]

7.3.5 active_subspace

- Keywords Area
Active (variable) subspace model

### Specification

**Alias:** subspace  
**Argument(s):** none  
**Child Keywords:**

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
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<td></td>
<td>Pointer to specify a full-space model, from which to construct a lower dimensional surrogate</td>
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<tr>
<td>Optional</td>
<td>initial_samples</td>
<td></td>
<td>Initial number of samples for sampling-based methods</td>
</tr>
<tr>
<td>Optional</td>
<td>sample_type</td>
<td></td>
<td>Selection of sampling strategy</td>
</tr>
<tr>
<td>Optional</td>
<td>truncation_method</td>
<td></td>
<td>Metric that estimates active subspace size</td>
</tr>
<tr>
<td>Optional</td>
<td>dimension</td>
<td></td>
<td>Explicitly specify the desired subspace size</td>
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<tr>
<td>Optional</td>
<td>bootstrap_samples</td>
<td></td>
<td>Number of bootstrap replicates used in truncation metrics</td>
</tr>
<tr>
<td>Optional</td>
<td>build_surrogate</td>
<td></td>
<td>Construct moving least squares surrogate over active subspace</td>
</tr>
</tbody>
</table>
Description

A model that transforms the original model (given by actual_model_pointer) to one with a reduced set of variables. This reduced model is identified by iteratively sampling the gradient of the original model and performing a singular value decomposition of the gradient matrix.

Expected Output

A subspace model will perform an initial sampling design to identify an active subspace using one of the truncation methods.

Usage Tips

If the desired subspace size is not identified, consider using the explicit dimension truncation option or one of the other truncation methods.

Examples

Perform an initial 100 gradient samples and use the bing_li truncation method to identify an active subspace. The truncation method uses 150 bootstrap samples to compute the Bing Li truncation metric.

model
subspace
  id_model = 'SUBSPACE'
  actual_model_pointer = 'FULLSPACE'
  initial_samples 100
  truncation_method bing_li
  bootstrap_samples 150

Theory

The idea behind active subspaces is to find directions in the input variable space in which the quantity of interest is nearly constant. After rotation of the input variables, this method can allow significant dimension reduction. Below is a brief summary of the process.

1. Compute the gradient of the quantity of interest, \( q = f(x) \), at several locations sampled from the full input space,
\[
\nabla_x f_i = \nabla f(x_i).
\]

2. Compute the eigendecomposition of the matrix \( \hat{C} \),
\[
\hat{C} = \frac{1}{M} \sum_{i=1}^{M} \nabla_x f_i \nabla_x f_i^T = \hat{W} \hat{\Sigma} \hat{W}^T,
\]
where \( \hat{W} \) has eigenvectors as columns, \( \hat{\Sigma} = \text{diag}(\hat{\lambda}_1, \ldots, \hat{\lambda}_N) \) contains eigenvalues, and \( N \) is the total number of parameters.

3. Using a truncation_method or specifying a dimension to estimate the active subspace size, split the eigenvectors into active and inactive directions,
\[
\hat{W} = [\hat{W}_1 \quad \hat{W}_2].
\]
These eigenvectors are used to rotate the input variables.

4. Next the input variables, \( x \), are expanded in terms of active and inactive variables,
\[
x = W_1 y + W_2 z.
\]
5. A surrogate is then built as a function of the active variables,

\[ g(y) \approx f(x) \]

For additional information, see:


7.3.5.1 actual_model_pointer

- Keywords Area
- model
- active_subspace
- actual_model_pointer

Pointer to specify a full-space model, from which to construct a lower dimensional surrogate

Topics

This keyword is related to the topics:

- block_pointer

Specification

Alias: none

Argument(s): STRING

Description

This must point to a model block, identified by id_model. That model will be run to generate gradient data, from which an active subspace model will be identified and built.

See block_pointer for details about pointers.

7.3.5.2 initial_samples

- Keywords Area
- model
- active_subspace
- initial_samples

Initial number of samples for sampling-based methods
Specification

Alias: none
Argument(s): INTEGER
Default: model-dependent

Description

The initial_samples keyword is used to define the number of initial samples (i.e., randomly chosen sets of variable values) at which to execute a model. The initial samples may later be augmented in an iterative process.

Default Behavior

By default, Dakota will use the minimum number of samples required by the chosen method.

Usage Tips

To obtain linear sensitivities or to construct a linear response surface, at least \(dim+1\) samples should be used, where "\(dim\)" is the number of variables. For sensitivities to quadratic terms or quadratic response surfaces, at least \((dim+1)(dim+2)/2\) samples are needed. For uncertainty quantification, we recommend at least \(10\times dim\) samples. For variance_based_decomp, we recommend hundreds to thousands of samples. Note that for variance_based_decomp, the number of simulations performed will be \(N\times (dim+2)\).

Examples

```plaintext
method
  sampling
    sample_type random
    initial_samples = 20
    refinement_samples = 5
```

7.3.5.3 sample_type

- Keywords Area
- model
- active_subspace
- sample_type

Selection of sampling strategy

Specification

Alias: none
Argument(s): none
Default: random
Child Keywords:
7.3 model

<table>
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<th>Dakota Keyword Description</th>
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<td>Uses Latin Hypercube Sampling (LHS) to sample variables</td>
</tr>
<tr>
<td></td>
<td></td>
<td>random</td>
<td>Uses purely random Monte Carlo sampling to sample variables</td>
</tr>
</tbody>
</table>

**Description**

The `sample_type` keyword allows the user to select between two types of sampling: Monte Carlo (pure random) and Latin hypercube (stratified) sampling.

The incremental keywords are deprecated; instead use `samples` together with `refinement_samples`.

**Default Behavior**

If the `sample_type` keyword is present, it must be accompanied by `lhs` or `random`. In most contexts, `lhs` is the default (exception: multilevel_sampling uses Monte Carlo by default).

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
    seed = 83921
```

### 7.3.5.3.1 lhs

- **Keywords Area**
- **model**
- **active_subspace**
- **sample_type**
- **lhs**

Uses Latin Hypercube Sampling (LHS) to sample variables

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

The `lhs` keyword invokes Latin Hypercube Sampling as the means of drawing samples of uncertain variables according to their probability distributions. This is a stratified, space-filling approach that selects variable values from a set of equi-probable bins.

**Default Behavior**
Latin Hypercube Sampling is the default sampling mode in most contexts (exception: multilevel_sampling). To explicitly specify LHS in the Dakota input file, the `lhs` keyword must appear in conjunction with the `sample_type` keyword.

**Usage Tips**

Latin Hypercube Sampling is very robust and can be applied to any problem. It is fairly effective at estimating the mean of model responses and linear correlations with a reasonably small number of samples relative to the number of variables.

**Examples**

```plaintext
method
  sampling
    sample_type lhs
    samples = 20
```

**7.3.5.3.2 random**

- **Keywords Area**
- **model**
- **active_subspace**
- **sample_type**
- **random**

Uses purely random Monte Carlo sampling to sample variables

**Specification**

- **Alias:** none
- **Argument(s):** none

**Description**

The `random` keyword invokes Monte Carlo sampling as the means of drawing samples of uncertain variables according to their probability distributions.

**Default Behavior**

In most contexts, Monte Carlo sampling is not the default sampling mode (exception: multilevel_sampling). To change this behavior, the `random` keyword must be specified in conjunction with the `sample_type` keyword.

**Usage Tips**

Monte Carlo sampling is more computationally expensive than Latin Hypercube Sampling as it requires a larger number of samples to accurately estimate statistics.

**Examples**

```plaintext
method
  sampling
    sample_type random
    samples = 200
```
7.3 model

7.3.5.4 truncation_method

- Keywords Area
- model
- active_subspace
- truncation_method

Metric that estimates active subspace size

Specification

Alias: none
Argument(s): none
Default: constantine
Child Keywords:

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<th>Description of Group</th>
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<td>Use the Bing Li &quot;ladle&quot; diagnostic to truncate subspace</td>
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<tr>
<td>Optional</td>
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<td>constantine</td>
<td>Use the Constantine diagnostic to truncate subspace</td>
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<td>Truncate the subspace based on eigenvalue energy</td>
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<tr>
<td>Optional</td>
<td></td>
<td>cross_validation</td>
<td>Truncate the subspace to minimize surrogate cross-validation error</td>
</tr>
</tbody>
</table>

Description

Metric that controls how many basis vectors are retained in the active subspace.

Default Behavior

The default is to use the constantine diagnostic.

Usage Tips

If the automated subspace identification methods do not yield desirable results, consider using the explicit dimension truncation option.

7.3.5.4.1 bing_li

- Keywords Area
- model
Use the Bing Li "ladle" diagnostic to truncate subspace

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Uses a trade-off criterion to determine where to truncate the active subspace. The criterion is a function of the eigenvalues and eigenvectors of the active subspace gradient matrix. This function compares the decrease in eigenvalue amplitude with the increase in eigenvector variability under bootstrap sampling of the gradient matrix. The active subspace size is taken to be the index of the first minimum of this quantity.

**Usage Tips**

If this automated diagnostic does not yield desirable results, consider using the explicit dimension truncation option or one of the other truncation methods.

**Theory**

Below is a brief outline of the Bing Li method of active subspace identification. The first two steps are common to all active subspace truncation methods.

1. Compute the gradient of the quantity of interest, \( q = f(x) \), at several locations sampled from the input space, \( \nabla_x f_i = \nabla f(x_i) \).

2. Compute the eigendecomposition of the matrix \( \hat{C} \),  
\[
\hat{C} = \frac{1}{M} \sum_{i=1}^{M} \nabla_x f_i \nabla_x f_i^T = \hat{W} \hat{\Lambda} \hat{W}^T,
\]

where \( \hat{W} \) has eigenvectors as columns, \( \hat{\Lambda} = \text{diag}(\hat{\lambda}_1, \ldots, \hat{\lambda}_N) \) contains eigenvalues, and \( N \) is the total number of parameters.

3. Normalize the eigenvalues,  
\[
\hat{\lambda}_i = \frac{\hat{\lambda}_i}{\sum_j \hat{\lambda}_j}.
\]

4. Use bootstrap sampling of the gradients found in step 1 to compute replicate eigendecompositions,  
\[
\hat{C}_j = \hat{W}_j \hat{\Lambda}_j \hat{W}_j^T.
\]

5. Compute variability of eigenvectors,  
\[
f^0_i = \frac{1}{M_{\text{boot}}} \sum_{j=1}^{M_{\text{boot}}} \left\{ 1 - | \det(\hat{W}_j^T \hat{W}_i^*) | \right\},
\]

where \( \hat{W}_i \) and \( \hat{W}_j^* \) both contain only the first \( i \) eigenvectors and \( M_{\text{boot}} \) is the number of bootstrap samples. The value of the variability at the first index, \( f^0_1 \), is defined as zero.
6. Normalize the eigenvector variability,

\[ f_i = \frac{f_i^0}{\sum_j f_j^0}. \]

7. The criterion, \( g_i \), is defined as,

\[ g_i = \lambda_i + f_i. \]

8. The index of first minimum of \( g_i \) is then the estimated active subspace rank.

For additional information, see Luo, Wei, and Bing Li. "Combining eigenvalues and variation of eigenvectors for order determination." SIAM, 2015.

7.3.5.4.2 constantine

- **Keywords**: Area
- **model**
- **active_subspace**
- **truncation_method**
- **constantine**

Use the Constantine diagnostic to truncate subspace

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Uses a criterion based on the variability of the subspace estimate. Eigenvectors are computed for bootstrap samples of the gradient matrix. The subspace size associated with the minimum distance between bootstrap eigenvectors and the nominal eigenvectors is the estimated active subspace size.

**Usage Tips**

If this automated diagnostic does not yield desirable results, consider using the explicit dimension truncation option or one of the other truncation methods.

**Theory**

Below is a brief outline of the Constantine method of active subspace identification. The first two steps are common to all active subspace truncation methods.

1. Compute the gradient of the quantity of interest, \( q = f(x) \), at several locations sampled from the input space,

\[ \nabla_x f_i = \nabla f(x_i). \]

2. Compute the eigendecomposition of the matrix \( \hat{C} \),

\[ \hat{C} = \frac{1}{M} \sum_{i=1}^{M} \nabla_x f_i \nabla_x f_i^T = \hat{W} \hat{\lambda} \hat{W}^T, \]

where \( \hat{W} \) has eigenvectors as columns, \( \hat{\lambda} = \text{diag}(\hat{\lambda}_1, \ldots, \hat{\lambda}_N) \) contains eigenvalues, and \( N \) is the total number of parameters.
3. Use bootstrap sampling of the gradients found in step 1 to compute replicate eigendecompositions,

\[ \tilde{C}_j = \tilde{W}_j \tilde{W}_j^T. \]

4. Compute the average distance between nominal and bootstrap subspaces,

\[ e_n^* = \frac{1}{M_{\text{boot}}} \sum_j \text{dist}(\text{ran}(\tilde{W}_n), \text{ran}(\tilde{W}_j,n)) = \frac{1}{M_{\text{boot}}} \sum_j \| \tilde{W}_n \tilde{W}_n^T - \tilde{W}_j,n \tilde{W}_j,n^T \|. \]

where \( M_{\text{boot}} \) is the number of bootstrap samples, \( \tilde{W}_n \) and \( \tilde{W}_j,n \) both contain only the first \( n \) eigenvectors, and \( n < N \).

5. The estimated subspace rank, \( r \), is then,

\[ r = \arg \min_ne_n^*. \]

For additional information, see Constantine, Paul G. "Active Subspaces: Emerging Ideas for Dimension Reduction in Parameter Studies". Vol. 2. SIAM, 2015.

7.3.5.4.3 energy

- Keywords Area
- model
- active_subspace
- truncation_method
- energy

Truncate the subspace based on eigenvalue energy

**Specification**

**Alias**: none

**Argument(s)**: none

**Child Keywords**:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tbody>
<tr>
<td>Optional</td>
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<td>truncation_- tolerance</td>
<td>Specify the maximum percentage (as a decimal) of the eigenvalue energy not captured by the active subspace representation.</td>
</tr>
</tbody>
</table>

**Description**

Uses a criterion based on the derivative matrix eigenvalue energy.

**Usage Tips**

This subspace truncation method may work best when working with non-normally distributed uncertain variables. If this automated diagnostic does not yield desirable results, consider using the explicit dimension truncation option or one of the other truncation methods.
Theory

Using the eigenvalue energy truncation metric, the subspace size is determined using the following equation:

\[ n = \inf \left\{ d \in \mathbb{Z} \mid 1 \leq d \leq N \land 1 - \frac{\sum_{i=1}^{d} \lambda_i}{\sum_{i=1}^{N} \lambda_i} < \varepsilon \right\} \]

where \( \varepsilon \) is the truncation_tolerance, \( n \) is the estimated subspace size, \( N \) is the size of the full space, and \( \lambda_i \) are the eigenvalues of the derivative matrix.

7.3.5.4.3.1 truncation_tolerance

- Keywords Area
- model
- active_subspace
- truncation_method
- energy
- truncation_tolerance

Specify the maximum percentage (as a decimal) of the eigenvalue energy not captured by the active subspace representation.

Specification

Alias: none
Argument(s): REAL

Description

The truncation_tolerance is used in the energy truncation method to identify an active subspace. The truncation tolerance is used in the following equation to determine the subspace size:

\[ n = \inf \left\{ d \in \mathbb{Z} \mid 1 \leq d \leq N \land 1 - \frac{\sum_{i=1}^{d} \lambda_i}{\sum_{i=1}^{N} \lambda_i} < \varepsilon \right\} \]

where \( \varepsilon \) is the truncation_tolerance, \( n \) is the estimated subspace size, \( N \) is the size of the full space, and \( \lambda_i \) are the eigenvalues of the derivative matrix.

The default value for truncation_tolerance is \( 1 \times 10^{-6} \).

7.3.5.4.4 cross_validation

- Keywords Area
- model
- active_subspace
- truncation_method
- cross_validation

Truncate the subspace to minimize surrogate cross-validation error.
Specification

**Alias:** none

**Argument(s):** none

**Default:** relative

**Child Keywords:**

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<td>exhaustive</td>
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**Dakota Keyword Description**
- Select subspace to minimize cross-validation error
- Choose subspace with cross-validation error less than tolerance
- Choose subspace where cross-validation error stabilizes
- Tolerance for cross-validation error value
- Tolerance for cross-validation error stabilization
- Maximum subspace dimension to consider
- Assess all admissible subspace dimensions

**Description**

Select the subspace dimension that minimizes the 10-fold cross-validation error when constructing a moving least squares surrogate model.

7.3.5.4.4.1 minimum

- Keywords Area
- model
- active_subspace
- truncation_method
7.3 model

- `cross_validation`
- `minimum`

Select subspace to minimize cross-validation error

**Specification**

*Alias*: none

*Argument(s)*: none

**Description**

Select the subspace dimension that minimizes the cross-validation error across sizes considered.

7.3.5.4.4.2 `relative`

- `Keywords Area`
- `model`
- `active_subspace`
- `truncation_method`
- `cross_validation`
- `relative`

Choose subspace with cross-validation error less than tolerance

**Specification**

*Alias*: none

*Argument(s)*: none

**Description**

Select smallest subspace with cross-validation error less than specified tolerance.

7.3.5.4.4.3 `decrease`

- `Keywords Area`
- `model`
- `active_subspace`
- `truncation_method`
- `cross_validation`
- `decrease`

Choose subspace where cross-validation error stabilizes
Specification

**Alias**: none

**Argument(s)**: none

Description

Select smallest subspace where change in cross-validation error to next larger subspace falls below specified tolerance.

7.3.5.4.4.4  **relative_tolerance**

- **Keywords Area**
- **model**
- **active_subspace**
- **truncation_method**
- **cross_validation**
- **relative_tolerance**

Tolerance for cross-validation error value

Specification

**Alias**: none

**Argument(s)**: REAL

**Default**: 1.0e-6

Description

Select smallest subspace where change in cross-validation error falls below specified tolerance.

7.3.5.4.4.5  **decrease_tolerance**

- **Keywords Area**
- **model**
- **active_subspace**
- **truncation_method**
- **cross_validation**
- **decrease_tolerance**

Tolerance for cross-validation error stabilization

Specification

**Alias**: none

**Argument(s)**: REAL

**Default**: 1.0e-6
Description

Select smallest subspace where change in cross-validation error falls below specified tolerance.

7.3.5.4.4.6 max_rank

- Keywords Area
- model
- active_subspace
- truncation_method
- cross_validation
- max_rank

Maximum subspace dimension to consider

Specification

Alias: none

Argument(s): INTEGER

Default: number fullspace vars

Description

In cross-validation, only consider subspace dimensions from 1 to max_rank.

7.3.5.4.4.7 exhaustive

- Keywords Area
- model
- active_subspace
- truncation_method
- cross_validation
- exhaustive

Assess all admissible subspace dimensions

Specification

Alias: none

Argument(s): none

Default: on

Description

Assess cross-validation error for all admissible subspace dimensions, even if a particular dimension meets specified cross-validation tolerance(s).
7.3.5.5 dimension

- Keywords Area
- model
- active_subspace
- dimension

Explicitly specify the desired subspace size

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

This control explicitly indicates the number of basis vectors to retain. The subspace model will include exactly `dimension` variables.

**Default Behavior**

Active for adapted basis. Not active for active subspace (the number of basis vectors will be chosen by one of the truncation methods).

**Usage Tips**

This control can be helpful when *a priori* studies give insight to the appropriate subspace size.

7.3.5.6 bootstrap_samples

- Keywords Area
- model
- active_subspace
- bootstrap_samples

Number of bootstrap replicates used in truncation metrics

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

The number of bootstrap replicates used to estimate the active subspace size.

**Default Behavior**

Use 100 replicates.
7.3.5.7 build_surrogate

- **Keywords Area**
- model
- active_subspace
- build_surrogate

Construct moving least squares surrogate over active subspace

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<tr>
<td>Optional</td>
<td></td>
<td>refinement_-samples</td>
<td>Number of supplementary surrogate build samples</td>
</tr>
</tbody>
</table>

**Description**

Once the active subspace variables have been identified, replace the fullspace truth model with a moving least squares surrogate model over the reduced variables. The surrogate is constructed using the fullspace function value samples collected during subspace identification, possibly supplemented with additional `refinement_samples` in the fullspace.

**Default Behavior**

No surrogate is constructed by default.

**Examples**

```dakota
model
active_subspace
  id_model = 'SUBSPACE'
  actual_model_pointer = 'FULLSPACE'
  initial_samples 100
build_surrogate
  refinement_samples 10
```

7.3.5.7.1 refinement_samples

- **Keywords Area**
- model
- active_subspace
- build_surrogate
- refinement_samples

Number of supplementary surrogate build samples
Specification

Alias: none
Argument(s): INTEGERLIST
Default: 0

Description

Augment the function value data collected during subspace identification with the specified number of refinement_samples to build the surrogate model.

7.3.5.8 normalization

- Keywords Area
- model
- active_subspace
- normalization

Normalize gradient samples

Specification

Alias: none
Argument(s): none

Child Keywords:

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<td>Normalize by sample mean of function values</td>
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<td>mean_gradient</td>
<td>Normalize by sample mean of gradient norms</td>
</tr>
<tr>
<td></td>
<td></td>
<td>local_gradient</td>
<td>Normalize each gradient sample by its norm</td>
</tr>
</tbody>
</table>

Description

Normalize the matrix of sampled gradients before identifying subspace.

Default Behavior

The default is local_gradient

7.3.5.8.1 mean_value

- Keywords Area
Normalize by sample mean of function values

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

For each response function $f_i$, normalize its gradient data by the mean $m_i$ of the function values taken across samples $j = 1, \ldots, J$:

$$m_i = \frac{1}{J} \sum_{j=1}^{J} f_i^j$$

7.3.5.8.2 **mean_gradient**

- **Keywords Area**
- **model**
- **active_subspace**
- **normalization**
- **mean_gradient**

Normalize by sample mean of gradient norms

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

For each response function $f_i$, normalize its gradient data by the mean $m_i$ of the gradient two-norms taken across samples $j = 1, \ldots, J$:

$$m_i = \frac{1}{J} \sum_{j=1}^{J} \| \nabla f_i^j \|_2$$
7.3.5.8.3 local_gradient

- Keywords Area
- model
- active_subspace
- normalization
- local_gradient

Normalize each gradient sample by its norm

Specification

Alias: none

Argument(s): none

Description

For each response function $f_i$, normalize each gradient sample $\nabla^j f_i$ by its two-norm $\|\nabla^j f_i\|

7.3.6 adapted_basis

- Keywords Area
- model
- adapted_basis

Basis adaptation model

Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
<td>Required</td>
<td>actual_model_pointer</td>
<td>truncation_tolerance</td>
<td>Pointer to specify a “truth” model, from which to construct a surrogate Convergence tolerance used to identify the (reduced) dimension of the rotation matrix</td>
</tr>
</tbody>
</table>
Description

A model that transforms the original model (given by actual_model_pointer) to one with a rotated set of variables. The current implementation does not support the reduction of the dimension for the new model and only transform the variable set in a rotated one in which the variables are arranged in a decreasing importance order.

An initial PCE representation is built with a sparse grid method and the PCE coefficients are then used to build a rotation matrix. This matrix is also reported as a result of the code execution. The new model can subsequently be used for an UQ workflow.

Expected Output  A basis adaptation model will perform an initial sparse grid design to identify the rotation matrix.

Examples

Perform an initial sparse grid design (level 1) to evaluate the PCE expansion and evaluate the rotation matrix. Afterwards, 100 samples are generated for the model in the rotated space to obtain its statistics.

```python
method,
    sampling
      model_pointer = 'SUBSPACE'
      samples = 100 seed = 1234567

model
  id_model = 'SUBSPACE'
  adapted_basis
    actual_model_pointer = 'FULLSPACE'
    sparse_grid_level = 1
```

Theory

The idea behind the Basis Adaptation method is to generate a PCE representation and rotate it such that the new set of rotated variables are organized in a decreasing importance ordering. Subsequently this rotation matrix can be truncated according to some criterion to only include the significant directions.

The first step of the Basis Adaptation is to compute a PCE expansion (we assume here $\xi$ to have a standard multivariate Gaussian distribution)

$$Q(\xi) = \sum_{\alpha \in J_{d,p}} Q_\alpha \psi_\alpha(\xi),$$

where $\alpha = (\alpha_1, \ldots, \alpha_d) \in J_{d,p} := (N_0)^d$ with $|\alpha| = \sum_{i=1}^d \alpha_i \leq d$ is multi-index of dimension $d$ and order up to $p$.

Afterwards, a multivariate Gaussian distribution $\eta$ is sought such that

$$\eta = A \xi,$$

where $A$ is an isometry such that $AA^\top = I$.

The basis adaptation model is obtained by expressing the original model with respect to the rotated set of variables $\eta$ as

$$Q^A(\eta) = \sum_{\beta \in J_{d,p}} Q^A_\beta \psi^A_\beta(\eta).$$

Since the basis for both the original and the adapted basis model span the same space, we know that $Q^A(\eta(\xi)) = Q(\xi)$ and therefore a relationships between the PCE coefficients exists

$$Q_\alpha = \sum_{\beta \in J_{d,p}} Q^A_\beta \langle \psi^A_\beta, \psi_\alpha \rangle, \ \alpha \in J_{d,p}. $$

The linear adaptation strategy is used at this time in Dakota to obtain the rotation matrix $A$. The steps to obtain the matrix $A$ are the following:
1. The first variable \( \eta_1 \) is obtained by directly combining the first order terms, that is

\[
\eta_1 = \sum_{\alpha \in J_d, 1} Q_\alpha \psi_\alpha(\xi) = \sum_{i=1}^{d} Q_i \xi_i,
\]

where \( e_i \) is the \( d \)-dimensional multi-index with 1 at the \( i \)th location and zero elsewhere.

2. The remaining rows of the \( A \) matrix (\( A \in \mathbb{R}^{d \times d} \)) are obtained by using the largest coefficient, row after row, placed in the column that it occupies in the first row. All other terms are set to zero.

3. As last step, in order to obtain an isometry, the Gram-Schmidt procedure is followed on \( A \).

See the theory manual for more details.

See Also

These keywords may also be of interest:

- active_subspace

7.3.6.1 actual_model_pointer

- Keywords Area
- model
- adapted_basis
- actual_model_pointer

Pointer to specify a "truth" model, from which to construct a surrogate

Topics

This keyword is related to the topics:

- block_pointer

Specification

**Alias:** none

**Argument(s):** STRING

**Child Keywords:**

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<tr>
<td>Required</td>
<td>sparse_grid_level</td>
<td>Level to use in sparse grid integration or interpolation</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Description

This must point to a model block, identified by `id_model`. That model will be run to generate training data, from which a surrogate model will be constructed.

See `block_pointer` for details about pointers.

#### 7.3.6.1.1 sparse_grid_level

- **Keywords** Area
- model
- adapted_basis
- actual_model_pointer
- sparse_grid_level

**Level to use in sparse grid integration or interpolation**

#### Specification

**Alias:** none

**Argument(s):** INTEGER

#### Description

Multi-dimensional integration by the Smolyak sparse grid method (specified with `sparse_grid_level` and, optionally, `dimension_preference`). The underlying one-dimensional integration rules are the same as for the tensor-product quadrature case; however, the default rule selection is nested for sparse grids (Genz-Keister for normals/transformed normals and Gauss-Patterson for uniforms/transformed uniforms). This default can be overridden with an explicit `non_nested` specification (resulting in Gauss-Hermite for normals/transformed normals and Gauss-Legendre for uniforms/transformed uniforms). As for tensor quadrature, the `dimension_preference` specification enables the use of anisotropic sparse grids (refer to the PCE description in the User's Manual for the anisotropic index set constraint definition). Similar to anisotropic tensor grids, the dimension with greatest preference will have resolution at the full `sparse_grid_level` and all other dimension resolutions will be reduced in proportion to their reduced preference. For PCE with either isotropic or anisotropic sparse grids, a summation of tensor-product expansions is used, where each anisotropic tensor-product quadrature rule underlying the sparse grid construction results in its own anisotropic tensor-product expansion as described in case 1. These anisotropic tensor-product expansions are summed into a sparse PCE using the standard Smolyak summation (again, refer to the User's Manual for additional details). As for quadrature_order, the `sparse_grid_level` specification admits an array input for enabling specification of multiple grid resolutions used by certain advanced solution methodologies.

A corresponding sequence specification is documented at, e.g., `sparse_grid_level_sequence` and `sparse_grid_level_sequence`
See Also

These keywords may also be of interest:

- `sparse_grid_level_sequence` method-multifidelity_stoch_collocation-sparse_grid_level_sequence

7.3.6.1.2 expansion_order

- Keywords Area
- model
- adapted_basis
- actual_model_pointer
- expansion_order

The (initial) order of a polynomial expansion

Specification

Alias: none

Argument(s): INTEGER

Child Keywords:

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<tbody>
<tr>
<td></td>
<td></td>
<td>collocation_ratio</td>
<td>Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.</td>
</tr>
</tbody>
</table>

Description

When the expansion_order for a polynomial chaos expansion is specified, the coefficients may be computed by integration based on random samples or by regression using either random or sub-sampled tensor product quadrature points.

Multidimensional integration by Latin hypercube sampling (specified with `expansion_samples`). In this case, the expansion order \( p \) cannot be inferred from the numerical integration specification and it is necessary to provide an expansion_order to specify \( p \) for a total-order expansion.

Linear regression (specified with either `collocation_points` or `collocation_ratio`). A total-order expansion is used and must be specified using `expansion_order` as described in the previous option. To avoid requiring the user to calculate \( N \) from \( n \) and \( p \), the `collocation_ratio` allows for specification of a constant factor applied to \( N \) (e.g., \( \text{collocation_ratio} = 2 \) produces samples = \( 2N \)). In addition, the default linear relationship with \( N \) can be overridden using a real-valued exponent specified using `ratio_order`. In this case, the number of samples becomes \( cN^o \) where \( c \) is the `collocation_ratio` and \( o \) is the `ratio_order`. The `use_derivatives` flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \( \frac{cN^o}{p+1} \)). When admissible, a constrained least squares approach is employed in which response values are first reproduced exactly and error in reproducing response derivatives is minimized. Two collocation grid options are supported: the default is Latin hypercube sampling.
("point collocation"), and an alternate approach of "probabilistic collocation" is also available through inclusion of the tensor_grid keyword. In this alternate case, the collocation grid is defined using a subset of tensor-product quadrature points: the order of the tensor-product grid is selected as one more than the expansion order in each dimension (to avoid sampling at roots of the basis polynomials) and then the tensor multi-index is uniformly sampled to generate a non-repeated subset of tensor quadrature points.

If collocation_points or collocation_ratio is specified, the PCE coefficients will be determined by regression. If no regression specification is provided, appropriate defaults are defined. Specifically SVD-based least-squares will be used for solving over-determined systems and under-determined systems will be solved using LASSO. For the situation when the number of function values is smaller than the number of terms in a PCE, but the total number of samples including gradient values is greater than the number of terms, the resulting over-determined system will be solved using equality constrained least squares. Technical information on the various methods listed below can be found in the Linear regression section of the Theory Manual. Some of the regression methods (OMP, LASSO, and LARS) are able to produce a set of possible PCE coefficient vectors (see the Linear regression section in the Theory Manual). If cross validation is inactive, then only one solution, consistent with the noise_tolerance, will be returned. If cross validation is active, Dakota will choose between possible coefficient vectors found internally by the regression method across the set of expansion orders (1, ..., expansion_order) and the set of specified noise tolerances and return the one with the lowest cross validation error indicator.

7.3.6.1.2.1 collocation_ratio

• Keywords Area
  • model
  • adapted_basis
  • actual_model_pointer
  • expansion_order
  • collocation_ratio

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion.

Specification

Alias: none

Argument(s): REAL

Description

Set the number of points used to build a PCE via regression to be proportional to the number of terms in the expansion. To avoid requiring the user to calculate N from n and p, the collocation_ratio allows for specification of a constant factor applied to N (e.g., collocation_ratio = 2, produces samples = 2N). In addition, the default linear relationship with N can be overridden using a real-valued exponent specified using ratio_order. In this case, the number of samples becomes \(cN^\alpha\) where \(c\) is the collocation_ratio and \(\alpha\) is the ratio_order. The use_derivatives flag informs the regression approach to include derivative matching equations (limited to gradients at present) in the least squares solutions, enabling the use of fewer collocation points for a given expansion order and dimension (number of points required becomes \(\frac{cN^\alpha}{n+\parallel}\)).

7.3.6.1.3 dimension

• Keywords Area
  • model
  • adapted_basis
Prescribe the number of active dimensions

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Description**

If a prescribed dimension for the rotation matrix is desired, this can be specified with the keyword `dimension`. In this case, the algorithm does not use the `truncation_method` or the `truncation_tolerance` to determine the truncation of the rotation matrix, but it just selects the first $d$ dimensions, where $d$ is assigned by `dimension`.

**Examples**

The following block

```model
    id_model = 'SUBSPACE'
    adapted_basis
        actual_model_pointer = 'FULLSPACE'
    sparse_grid_level = 1
    dimension = 3
```

prescribes the selection of the first three most important directions to build the rotation matrix and the reduced model representation.

### 7.3.6.1.4 rotation_method

- **Keywords Area**
- **model**
- **adapted_basis**
- **actual_model_pointer**
- **rotation_method**

Method used to build the rotation matrix

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** norm

**Child Keywords:**
Description

The rotation matrix for the Adapted Basis method is built starting from a matrix which includes the linear PCE coefficients. The entries of the first row are the linear PCE coefficients. For all the other rows, each row has only one nonzero entry. Depending on the way used to construct all the rows, from the second to last one, there are two implemented methods, unranked and ranked. A followed Gram-schmidt process is applied on the resulting matrix to make it an isometry.

Default Behavior

The default is to use the ranked.

Usage Tips

Examples

The following method block

```dakota
model
  id_model = 'SUBSPACE'
  adapted_basis
    actual_model_pointer = 'FULLSPACE'
    sparse_grid_level = 1
    rotation_method = unranked
```  
changes the default method to the unranked method.

7.3.6.1.4.1  unranked

- Keywords Area
- model
- adapted_basis
- actual_model_pointer
- rotation_method
- unranked

Use the unranked method to obtain the rotation matrix

Specification

Alias: none

Argument(s): none
Description

In this case the construction of the second to last rows of the rotation matrix is performed by using the value 1 in the diagonal terms and zeros elsewhere.

7.3.6.1.4.2 ranked

- Keywords Area
- model
- adapted_basis
- actual_model_pointer
- rotation_method
- ranked

Use the ranked method to obtain the rotation matrix

Specification

Alias: none

Argument(s): none

Description

In this case the construction of the $i$th row, with $i \geq 2$, of the rotation matrix is performed by putting the $(i - 1)$th largest linear PCE coefficient in the column that it occupies in the first row.

Theory

The first row of the rotation captures the complete Gaussian components and is suggested as the leading direction. By the construction in this method, the other rotated variables are the variables in the original space that have the greatest sensitivity in descending order. The particular construction makes sure that the importance of the adapted variables is in descending order, which permits the possibility of reduced representation.

7.3.6.2 truncation_tolerance

- Keywords Area
- model
- adapted_basis
- truncation_tolerance

Convergence tolerance used to identify the (reduced) dimension of the rotation matrix

Specification

Alias: none

Argument(s): REAL

Default: 0.8
7.3 model

Description

This convergence tolerance controls the dimension of the rotation matrix after the truncation. The metric controlled by this convergence tolerance is a quantity varying between \([0,1]\). If a more aggressive dimension truncation is desired, the default value (0.8) can be reduced to a lower value.

Examples

The following block

```plaintext
model
  id_model = 'SUBSPACE'
  adapted_basis
    actual_model_pointer = 'FULLSPACE'
  sparse_grid_level = 1
  truncation_tolerance = 0.9
```

increases the default value (0.8) for the `truncation_tolerance` to 0.9 to obtain a rotation matrix and, consequently, a reduced model that more closely approximates the full dimensional model.

7.3.7 random_field

• Keywords Area
• model
• random_field

Experimental capability to generate a random field representation. from data, from simulation runs, or from a covariance matrix. The representation may then be sampled for use as a random field input to another simulation. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none
Argument(s): none

Child Keywords:

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<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>build_source</td>
<td>Specify how the random field will be built: from a data file, from simulation runs, or from a covariance matrix. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
</tbody>
</table>
### Description

Capability to generate a random field representation from data, from simulation runs, or from a covariance matrix. The random field may then be sampled for use as a random field input to another simulation. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

**Default Behavior** There are three main sections of the `random_field` model. The first section tells Dakota what data to use to build the random field. This is specified with `build_source`. The source of data to build
the random field may be a file with data (where the N rows of data correspond to N samples of the random field and the M columns correspond to field values), or it may be a simulation that generates field data, or it may be specified given a mesh and a covariance matrix governing how the field varies over the mesh. In the case of using a simulation to generate field data, the simulation is defined with `dace_method_pointer`. In the case of using a mesh and a covariance, the form of the covariance is defined with `analytic_covariance`.

The next section of the random field model specifies the form of the expansion, `expansion_form`. This can be either a Karhunen-Loeve expansion or a Principal components analysis. These are very similar: both involve the eigenvalues of the covariance matrix of the field data. The only difference is in the treatment of the estimation of the coefficients of the eigenvector basis functions. In the PCA case, we have developed an approach which makes the coefficients explicit functions of the uncertain variables used to generate the random field. The specification of the random field can also include the number of bases to retain or a truncation tolerance, which defines the percent variance that the expansion should capture.

The final section of the random field model allows the user to specify a pointer to a model over which the random field will be propagated, meaning the model which will be driven with the random field input. This part of the specification is optional: one can build a random field but not use it in a downstream model.

**Examples**

As stated above, this is an emerging capability. The syntax currently looks like the following:

```plaintext
random_field
  build_source
    rf_data_file | dace_method_pointer | analytic_covariance
  expansion_form
    karhunen_loeve | principal_components
  expansion_bases
  truncation_tolerance
  propagation_model_pointer
```

### 7.3.7.1 build_source

- **Keywords Area**

- **model**

- **random_field**

- **build_source**

Specify how the random field will be built: from a data file, from simulation runs, or from a covariance matrix. **THIS IS AN EXPERIMENTAL CAPABILITY.**

**Specification**

- **Alias:** none
- **Argument(s):** none
- **Child Keywords:**
### Description

As part of the capability to generate a random field representation, the user needs to specify the data used to generate the random field representation. This data may reside in a data file, it may be generated by running a set of simulations and generating field responses, or it may be generated by a covariance matrix defined over a mesh. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

**Default Behavior** Currently, the `build_source` that is fully working is the `dace_method_pointer`. The others are not fully operational.

#### 7.3.7.1.1 rf_data_file

- Keywords Area
  - model
  - random_field
  - build_source
  - rf_data_file

Specify that the random field will be built from a file of data. THIS IS AN EXPERIMENTAL CAPABILITY.

### Specification

**Alias:** none
7.3 model

Argument(s): STRING

Default: none

Description

As part of the capability to generate a random field representation, the user needs to specify the data used to generate the random field representation. The rows of this file represent separate samples, and the columns represent the field data. For example, if you have 100 samples of a random field of length 500, the data file will be of dimension 100 x 500. Currently, this option is not operational. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  Currenty, the build_source that is fully working is the dace_method_pointer. The others are not fully operational.

7.3.7.1.2 dace_method_pointer

- Keywords Area
- model
- random_field
- build_source
- dace_method_pointer

Pointer to a DACE method for purposes of generating an ensemble of field responses to be used in estimating a random field model. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none

Argument(s): STRING

Default: no design of experiments data

Description

As part of the capability to generate a random field representation, the user needs to specify the data used to generate the random field representation. One way to do this is to run a set of simulations and generating field responses. Dakota will then take the full set of field responses (e.g. multiple samples, where each sample has a field response) and construct a random field model representing the uncertainty in the ensemble. The dace_method_pointer is a pointer to a Design and Analysis of Computer Experiments (DACE) method, typically which is a sampling method. In this case, the sampling method should be on a simulation which can generate field responses (e.g. field_responses, field_objectives, or field_calibration terms.)

THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  Currently, the build_source that is fully working is the dace_method_pointer. The others are not fully operational.

7.3.7.1.3 analytic_c covariance

- Keywords Area
- model
- random_field
- build_source
• **analytic_covariance**

Use an analytic covariance function for the purposes of generating a random field model. THIS IS AN EXPERIMENTAL CAPABILITY.

**Specification**

**Alias:** none  
**Argument(s):** none  

**Child Keywords:**

<table>
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<tr>
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</tr>
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<tbody>
<tr>
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<td></td>
<td>squared_exponential</td>
<td>Specify a squared exponential covariance in the case where the random field is built from an analytic covariance function. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>exponential</td>
<td>Specify an exponential covariance in the case where the random field is built from an analytic covariance function. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
</tbody>
</table>

**Description**

As part of the capability to generate a random field representation, the user needs to specify the data used to generate the random field representation. If **analytic_covariance** is specified, an analytic covariance function will be used to generate instantiations of a random field over a mesh. The form of the covariance function must be specified (e.g. exponential or squared_exponential). THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

**Default Behavior**  Currently, the **build_source** that is fully working is the **dace_method_pointer**. The others are not fully operational.

7.3.7.1.3.1 squared_exponential

• **Keywords Area**
  
• **model**
  
• **random_field**
  
• **build_source**
  
• **analytic_covariance**
7.3 model

• squared_exponential

Specify a squared exponential covariance in the case where the random field is built from an analytic covariance function. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none
Argument(s): none

Description

THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  Currently, the build_source that is fully working is the dace_method_pointer. The others are not fully operational.

7.3.7.1.3.2 exponential

• Keywords Area
• model
• random_field
• build_source
• analytic_covariance
• exponential

Specify an exponential covariance in the case where the random field is built from an analytic covariance function. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none
Argument(s): none

Description

THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  Currently, the build_source that is fully working is the dace_method_pointer. The others are not fully operational.

7.3.7.2 expansion_form

• Keywords Area
• model
• random_field
• expansion_form

Specify the form of the expansion to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.
Specification

Alias: none

Argument(s): none

Child Keywords:

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<tr>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>karhunen_loeve</td>
<td>Specify Karhunen-Loeve as the expansion form to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>principal_components</td>
<td>Specify Principal Components as the form of the expansion to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.</td>
</tr>
</tbody>
</table>

Description

This control allows the user to specify the form of the random field representation (e.g. either a Karhunen-Loeve or Principal Components expansion). THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  The default expansion form is Karhunen-Loeve.

7.3.7.2.1 karhunen_loeve

- Keywords Area
- model
- random_field
- expansion_form
- karhunen_loeve

Specify Karhunen-Loeve as the expansion form to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none

Argument(s): none
Description

THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior The default expansion form is Karhunen-Loeve.

7.3.7.2.2 principal_components

• Keywords Area
• model
• random_field
• expansion_form
• principal_components

Specify Principal Components as the form of the expansion to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none
Argument(s): none

Description

Principal Components and Karhunen-Loeve are very similar functional forms of the expansion. They both use basis functions which are eigenvectors of the covariance matrix of the random field data. However, in principal components, we parameterize the coefficients of the expansion to be Gaussian process models which are functions of the uncertain parameters used to generate the initial ensemble of data representing the random field. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior The default expansion form is Karhunen-Loeve.

7.3.7.3 expansion_bases

• Keywords Area
• model
• random_field
• expansion_bases

Specify the number of basis functions to be used in the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none
Argument(s): INTEGER
Description

This control allows the user to specify the number of basis functions to be used in the random field representation (e.g. either a Karhunen-Loeve or Principal Components expansion). Typically, a small number of basis functions (3-5) will be sufficient to represent a significant amount of the variance in the response field. However, this depends on the particulars of the problem. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  If the user specifies neither truncation_tolerance nor expansion_bases, then the number of expansion bases that captures 95% of the variance will be used.

7.3.7.4 truncation_tolerance

- Keywords Area
- model
- random_field
- truncation_tolerance

Specify a percent of the response variance that should be captured with the random field representation. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none

Argument(s): REAL

Description

This control allows the user to specify the percent variance to be capture in the random field representation (e.g. either a Karhunen-Loeve or Principal Components expansion). Typically, the user would specify something like 0.9 or 0.95, where 0.9 means that 90% of the variance of the response field will be captured by the random field. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior  If the user specifies neither truncation_tolerance nor expansion_bases, then the truncation tolerance is set to 0.95 and the number of expansion bases that captures 95% of the variance will be used.

7.3.7.5 propagation_model_pointer

- Keywords Area
- model
- random_field
- propagation_model_pointer

Pointer to the model that will accept realizations of the random field and use them for subsequent analysis. Typically, this model will take the random field as inputs, e.g. a random field defining a pressure boundary or temperature boundary condition over a structure. THIS IS AN EXPERIMENTAL CAPABILITY.

Specification

Alias: none

Argument(s): STRING
Description

After a random field representation has been generated, Dakota will generate samples of that random field based on the representation. These sample realizations can then be used to drive another set of simulation analyses. The `propagation_model_pointer` is the model on which the random field realizations will be propagated. THIS IS AN EXPERIMENTAL CAPABILITY UNDER ACTIVE DEVELOPMENT.

Default Behavior

7.3.8 variables_pointer

- **Keywords Area**
- `model`
- `variables_pointer`

Specify which variables block will be included with this model block

Topics

This keyword is related to the topics:

- `block_pointer`

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** model use of last variables parsed

Description

The `variables_pointer` is used to specify which variables block will be used by the model, by cross-referencing with `id_variables` keyword in the `variables` block.

See `block_pointer` for details about pointers.

Default Behavior

When a variables pointer is not specified, the model will use the last variables block parsed from the input file.

7.3.9 responses_pointer

- **Keywords Area**
- `model`
- `responses_pointer`

Specify which responses block will be used by this model block

Topics

This keyword is related to the topics:

- `block_pointer`
Specification

**Alias:** none  

**Argument(s):** STRING  

**Default:** model use of last responses parsed

**Description**

The `responses_pointer` is used to specify which responses block will be used by the model, by cross-referencing with `id_responses` keyword in the `responses` block.  

See `block_pointer` for details about pointers.

**Default Behavior**

When a responses pointer is not specified, the model will use the last responses block parsed from the input file.

7.3.10 hierarchical_tagging

- **Keywords Area**
- **model**
- **hierarchical_tagging**

Enables hierarchical evaluation tagging

**Specification**

**Alias:** none  

**Argument(s):** none  

**Default:** no hierarchical tagging

**Description**

The hierarchical tagging option is useful for studies involving multiple models with a nested or hierarchical relationship. For example a nested model has a sub-method, which itself likely operates on a sub-model, or a hierarchical approximation involves coordination of low and high fidelity models. Specifying `hierarchical_tagging` will yield function evaluation identifiers (“tags”) composed of the evaluation IDs of the models involved, e.g., `outermodel.-innermodel.interfaceid = 4.9.2`. This communicates the outer contexts to the analysis driver when performing a function evaluation.

**Examples**

`test/dakota_uq_timeseries_ivp_optinterf.in` `test/dakota_uq_timeseries_sop_optinterf.in`

**See Also**

These keywords may also be of interest:

- `file_tag` model-nested
7.4 variables

- Keywords Area
- variables

Specifies the parameter set to be iterated by a particular method.

Topics

This keyword is related to the topics:

- block

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<td>id_variables</td>
<td>Name the variables block; helpful when there are multiple</td>
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<td>Optional</td>
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<td>active</td>
<td>Set the active variables view a method will see</td>
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<td>Optional/Choose One</td>
<td>Variable Domain (Group 1)</td>
<td>mixed</td>
<td>Maintain continuous/discrete variable distinction</td>
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<td></td>
<td></td>
<td>relaxed</td>
<td>Allow treatment of discrete variables as continuous</td>
</tr>
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<td>Design variable - continuous</td>
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<td></td>
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<td>Design variable - discrete range-valued</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>discrete_design_-set</td>
<td>Design variable - discrete set-valued</td>
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<td>Aleatory uncertain variable - normal (Gaussian)</td>
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<td>Aleatory uncertain discrete variable - binomial</td>
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<td>Aleatory uncertain discrete variable - geometric</td>
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<td>Correlation among aleatory uncertain variables</td>
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<td>------------------------------</td>
<td>------------------------------------------------</td>
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<td>Epistemic uncertain variable - values from one or more continuous intervals</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>discrete_interval_uncertain</td>
<td>Epistemic uncertain variable - values from one or more discrete intervals</td>
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</tr>
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<td>Optional</td>
<td>discrete_uncertain_set</td>
<td>Epistemic uncertain variable - discrete set-valued</td>
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<td>State variable - continuous</td>
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<td>discrete_state_range</td>
<td>State variables - discrete range-valued</td>
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<tr>
<td>Optional</td>
<td>discrete_state_set</td>
<td>State variable - discrete set-valued</td>
<td></td>
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<td>Define coefficients of the linear inequality constraints</td>
<td></td>
</tr>
<tr>
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<td>Define lower bounds for the linear inequality constraint</td>
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<tr>
<td>Optional</td>
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<td>Define upper bounds for the linear inequality constraint</td>
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<td>linear_inequality_scale_types</td>
<td>How to scale each linear inequality constraint</td>
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<tr>
<td>Optional</td>
<td>linear_inequality_scales</td>
<td>Characteristic values to scale linear inequalities</td>
<td></td>
</tr>
</tbody>
</table>
Description

The variables specification in a Dakota input file specifies the parameter set to be iterated by a particular method. In the case of:

- An optimization study: These variables are adjusted in order to locate an optimal design.
- Parameter studies/sensitivity analysis/design of experiments: These parameters are perturbed to explore the parameter space.
- Uncertainty analysis: The variables are associated with distribution/interval characterizations which are used to compute corresponding distribution/interval characterizations for response functions.

To accommodate these different studies, Dakota supports different:

- Variable types
  - design
  - aleatory uncertain
  - epistemic uncertain
  - state
- Variable domains
  - continuous
  - discrete
    - discrete range
    - discrete integer set
    - discrete string set
    - discrete real set

Use the variables page to browse the available variables by type and domain.

Variable Types

- Design Variables
  - Design variables are those variables which are modified for the purposes of seeking an optimal design.
  - The most common type of design variables encountered in engineering applications are of the continuous type. These variables may assume any real value within their bounds.
  - All but a handful of the optimization algorithms in Dakota support continuous design variables exclusively.
7.4 variables

- Aleatory Uncertain Variables
  - Aleatory uncertainty is also known as inherent variability, irreducible uncertainty, or randomness.
  - Aleatory uncertainty is predominantly characterized using probability theory. This is the only option implemented in Dakota.

- Epistemic Uncertain Variables
  - Epistemic uncertainty is uncertainty due to lack of knowledge.
  - In Dakota, epistemic uncertainty is assessed by interval analysis or the Dempster-Shafer theory of evidence.
  - Continuous or discrete interval or set-valued variables are used to define set-valued probabilities or basic probability assignments (BPA) which define a belief structure.
  - Note that epistemic uncertainty can also be modeled with probability density functions (as done with aleatory uncertainty). Dakota does not support this capability.

- State Variables
  - State variables consist of “other” variables which are to be mapped through the simulation interface, in that they are not to be used for design and they are not modeled as being uncertain.
  - State variables provide a convenient mechanism for managing additional model parameterizations such as mesh density, simulation convergence tolerances, and time step controls.
  - Only parameter studies and design of experiments methods will iterate on state variables.
  - The initial_value is used as the only value for the state variable for all other methods, unless active state is invoked.
  - See more details on the state_variables page.

Variable Domains
Continuous variables are typically defined by bounds. Discrete variables can be defined in one of three ways, which are discussed on the page discrete_variables.

Ordering of Variables
The ordering of variables is important, and a consistent ordering is employed throughout the Dakota software. The ordering is shown in dakota.input.summary (and in the hierarchical order of this reference manual) and can be summarized as:

1. design
   (a) continuous
   (b) discrete integer
   (c) discrete string
   (d) discrete real

2. aleatory uncertain
   (a) continuous
   (b) discrete integer
   (c) discrete string
   (d) discrete real

3. epistemic uncertain
   (a) continuous
   (b) discrete integer
   (c) discrete string
   (d) discrete real
4. state
  (a) continuous
  (b) discrete integer
  (c) discrete string
  (d) discrete real

Ordering of variable types below this granularity (e.g., from normal to histogram bin within aleatory uncertain - continuous) is defined somewhat arbitrarily, but is enforced consistently throughout the code.

**Active Variables**

The reason variable types exist is that methods have the capability to treat variable types differently. All methods have default behavior that determines which variable types are "active" and will be assigned values by the method. For example, optimization methods will only vary the design variables - by default.

The default behavior should be described on each method page, or on topics pages that relate to classes of methods. In addition, the default behavior can be modified using the active keyword.

At least one type of variables that are active for the method in use must have nonzero size (at least 1 active variable) or an input error message will result.

**Inferred Default Values and Bounds**

The concept of active variables allows any Dakota variable type to be used in any method context. Some methods, e.g., bound-constrained optimization or multi-dimensional or centered parameter studies, require bounds and/or an initial point on the variables, however uncertain variables may not be naturally defined in terms of these characteristics.

Distribution lower and upper bounds are explicit portions of the normal, lognormal, uniform, loguniform, triangular, and beta specifications, whereas they are implicitly defined for others. For example, bounds are naturally defined for histogram bin, histogram point, and interval variables (from the extreme values within the bin/point/interval specifications) as well as for binomial (0 to num_trials) and hypergeometric (0 to min(num_drawn, num_selected)) variables.

If not specified, distribution bounds are also inferred for normal and lognormal (if optional bounds are unspecified) as well as for exponential, gamma, gumbel, frechet, weibull, poisson, negative binomial, and geometric (which have no bounds specifications); these bounds are \([0, \mu + 3\sigma]\) for exponential, gamma, frechet, weibull, poisson, negative binomial, geometric, and unspecified lognormal, and \([\mu - 3\sigma, \mu + 3\sigma]\) for gumbel and unspecified normal.

When an initial point is needed and not explicitly specified in user input, it is assigned as described in the initial_point or initial_state specification, e.g., initial_point. For example, uncertain variables are initialized to their means, where mean values for bounded normal and bounded lognormal may be further adjusted to satisfy any user-specified distribution bounds, mean values for discrete integer range distributions are rounded down to the nearest integer, and mean values for discrete set distributions are rounded to the nearest set value.

**Examples**

Several examples follow. In the first example, two continuous design variables are specified:

```
variables,
  continuous_design = 2
  initial_point  0.9  1.1
  upper_bounds  5.8  2.9
  lower_bounds  0.5  -2.9
  descriptors 'radius' 'location'
```

In the next example, defaults are employed. In this case, initial_point will default to a vector of 0. values, upper_bounds will default to vector values of DBL_MAX (the maximum number representable in double precision for a particular platform), lower_bounds will default to a vector of -DBL_MAX values, and descriptors will default to a vector of 'cdv_i' strings, where i ranges from one to two:
7.4 variables

In the following example, the syntax for a normal-lognormal distribution is shown. One normal and one lognormal uncertain variable are completely specified by their means and standard deviations. In addition, the dependence structure between the two variables is specified using the uncertain_correlation_matrix.

variables,
    normal_uncertain = 1
    means = 1.0
    std_deviations = 1.0
    descriptors = 'TF1n'
    lognormal_uncertain = 1
    means = 2.0
    std_deviations = 0.5
    descriptors = 'TF2ln'
    uncertain_correlation_matrix = 1.0 0.2
                                 0.2 1.0

An example of the syntax for a state variables specification follows:

variables,
    continuous_state = 1
    initial_state 4.0
    lower_bounds 0.0
    upper_bounds 8.0
    descriptors 'CS1'
    discrete_state_range = 1
    initial_state 104
    lower_bounds 100
    upper_bounds 110
    descriptors 'DS1'

And in a more advanced example, a variables specification containing a set identifier, continuous and discrete design variables, normal and uniform uncertain variables, and continuous and discrete state variables is shown:

variables,
    id_variables = 'V1'
    continuous_design = 2
    initial_point 0.9 1.1
    upper_bounds 5.8 2.9
    lower_bounds 0.5 -2.9
    descriptors 'radius' 'location'
    discrete_design_range = 1
    initial_point 2
    upper_bounds 1
    lower_bounds 3
    descriptors 'material'
    normal_uncertain = 2
    means = 248.89, 593.33
    std_deviations = 12.4, 29.7
    descriptors = 'TF1n' 'TF2n'
    uniform_uncertain = 2
    lower_bounds = 199.3, 474.63
    upper_bounds = 298.5, 712.
    descriptors = 'TF1u' 'TF2u'
    continuous_state = 2
    initial_state = 1.e-4 1.e-6
    descriptors = 'EPSIT1' 'EPSIT2'
    discrete_state_set
        integer = 1
        initial_state = 100
        set_values = 100 212 375
        descriptors = 'load_case'

7.4.1 id_variables

- Keywords Area
• variables
  • id_variables

Name the variables block; helpful when there are multiple

Topics

This keyword is related to the topics:

• block_identifier

Specification

Alias: none
Argument(s): STRING
Default: use of last variables parsed

Description

The optional id_variables keyword accepts a string that uniquely identifies this variables block. A model can then use these variables by specifying the same string in its variables_pointer specification.

Default Behavior

If the id_variables specification is omitted, a particular variables specification will be used by a model only if that model does not include an variables_pointer and the variables block was the last (or only) one parsed.

Usage Tips

• It is a best practice to always use explicit variables IDs and pointers to avoid confusion.
• If only one variables block exists, then id_variables can be safely omitted from the variables block (and variables_pointer omitted from the model specification(s)), since there is no ambiguity.

Examples

For example, a model specification including

```model
  variables_pointer = 'V1'
```

will link to a response set with

```id_variables = 'V1'
```

7.4.2 active

• Keywords Area
  • variables
  • active

Set the active variables view a method will see
7.4 variables

Specification

**Alias:** none

**Argument(s):** none

**Default:** Infer from response or method specification

**Child Keywords:**

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<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Active Variables (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</tr>
<tr>
<td></td>
<td></td>
<td>aleatory</td>
<td>Option for the active keyword</td>
</tr>
<tr>
<td></td>
<td></td>
<td>epistemic</td>
<td>Option for the active keyword</td>
</tr>
<tr>
<td></td>
<td></td>
<td>state</td>
<td>Option for the active keyword</td>
</tr>
</tbody>
</table>

**Description**

There are certain situations where the user may want to explicitly control the subset of variables that is considered active for a certain Dakota method. This is done by specifying the keyword `active` in the variables specification block, followed by one of the following: `all`, `design`, `uncertain`, `aleatory`, `epistemic`, or `state`.

Specifying one of these subsets of variables will allow the Dakota method to operate on the specified variable types and override the default active subset.

If the user does not specify any explicit override of the active view of the variables, Dakota first considers the response function specification.

- If the user specifies objective functions or calibration terms in the response specification block, then we can infer that the active variables should be the design variables (since design variables are used within optimization and least squares methods).

- If the user instead specifies the generic response type of `response_functions`, then Dakota cannot infer the active variable subset from the response specification and will instead infer it from the method selection.

  1. If the method is a parameter study, or any of the methods available under dace, psuade, or fsu methods, the active view is set to all variables.

  2. For uncertainty quantification methods, if the method is sampling, then the view is set to aleatory if only aleatory variables are present, epistemic if only epistemic variables are present, or uncertain (covering both aleatory and epistemic) if both are present.

  3. If the uncertainty method involves interval estimation or evidence calculations, the view is set to epistemic.

  4. For other uncertainty quantification methods not mentioned in the previous sentences (e.g., reliability methods or stochastic expansion methods), the default view is set to aleatory.

  5. Finally, for verification studies using the Richardson extrapolation method, the active view is set to state.

  6. Note that in surrogate-based optimization, where the surrogate is built on points defined by the method defined by the `dace_method_pointer`, the sampling used to generate the points is performed only over the design variables as a default unless otherwise specified (e.g. state variables will not be sampled for surrogate construction).
As alluded to in the previous section, the iterative method selected for use in Dakota determines what subset, or view, of the variables data is active in the iteration. The general case of having a mixture of various different types of variables is supported within all of the Dakota methods even though certain methods will only modify certain types of variables (e.g., optimizers and least squares methods only modify design variables, and uncertainty quantification methods typically only utilize uncertain variables). This implies that variables which are not under the direct control of a particular iterator will be mapped through the interface in an unmodified state. This allows for a variety of parameterizations within the model in addition to those which are being used by a particular iterator, which can provide the convenience of consolidating the control over various modeling parameters in a single file (the Dakota input file). An important related point is that the variable set that is active with a particular iterator is the same variable set for which derivatives are typically computed.

Examples

For example, the default behavior for a nondeterministic sampling method is to sample the uncertain variables. However, if the user specified active all in the variables specification block, the sampling would be performed over all variables (e.g. design and state variables in addition to the uncertain variables). This may be desired in situations such as surrogate-based optimization under uncertainty, where a surrogate may be constructed to span both design and uncertain variables. This is an example where we expand the active subset beyond the default, but in other situations, we may wish to restrict from the default. An example of this would be performing design of experiments in the presence of multiple variable types (for which all types are active by default), but only wanting to sample over the design variables for purposes of constructing a surrogate model for optimization.

Theory

The optional status of the different variable type specifications allows the user to specify only those variables which are present (rather than explicitly specifying that the number of a particular type of variables is zero). However, at least one type of variables that are active for the iterator in use must have nonzero size or an input error message will result.

### 7.4.2.1 all

- Keywords Area
- variables
- active
- all

Option for the active keyword

Specification

**Alias:** none

**Argument(s):** none

**Description**

See the active keyword

### 7.4.2.2 design

- Keywords Area
7.4 variables

- variables
- active
- design

Option for the active keyword

Specification

Alias: none
Argument(s): none

Description

See the active keyword

7.4.2.3 uncertain

- Keywords Area
- variables
- active
- uncertain

Option for the active keyword

Specification

Alias: none
Argument(s): none

Description

See the active keyword

7.4.2.4 aleatory

- Keywords Area
- variables
- active
- aleatory

Option for the active keyword

Specification

Alias: none
Argument(s): none
Description

See the active keyword

7.4.2.5 epistemic

- Keywords Area
- variables
- active
- epistemic

Option for the active keyword

Specification

Alias: none

Argument(s): none

Description

See the active keyword

7.4.2.6 state

- Keywords Area
- variables
- active
- state

Option for the active keyword

Specification

Alias: none

Argument(s): none

Description

See the active keyword

7.4.3 mixed

- Keywords Area
- variables
- mixed

Maintain continuous/discrete variable distinction
7.4 variables

Specification

Alias: none
Argument(s): none
Default: relaxed (branch and bound), mixed (all other methods)

Description

The variables domain specifies how the discrete variables are treated. If the user specifies mixed in the variable specification block, the continuous and discrete variables are treated separately. If the user specifies relaxed in the variable specification block, the discrete variables are relaxed and treated as continuous variables. This may be useful in optimization problems involving both continuous and discrete variables when a user would like to use an optimization method that is designed for continuous variable optimization. All Dakota methods have a default value of mixed for the domain type except for the branch-and-bound method which has a default domain type of relaxed. Note that the branch-and-bound method is under development at this time. Finally, note that the domain selection applies to all variable types: design, aleatory uncertain, epistemic uncertain, and state.

With respect to domain type, if the user does not specify an explicit override of mixed or relaxed, Dakota infers the domain type from the method. As mentioned above, all methods currently use a mixed domain as a default, except the branch-and-bound method which is under development.

See Also

These keywords may also be of interest:

- relaxed

7.4.4 relaxed

- Keywords Area
- variables
- relaxed

Allow treatment of discrete variables as continuous

Specification

Alias: none
Argument(s): none

Description

The variables domain specifies how the discrete variables are treated. If the user specifies mixed in the variable specification block, the continuous and discrete variables are treated separately. If the user specifies relaxed in the variable specification block, the discrete variables are relaxed and treated as continuous variables. This may be useful in optimization problems involving both continuous and discrete variables when a user would like to use an optimization method that is designed for continuous variable optimization. All Dakota methods have a default value of mixed for the domain type except for the branch-and-bound method which has a default domain type of relaxed. Note that the branch-and-bound method is under development at this time. Finally, note that the domain selection applies to all variable types: design, aleatory uncertain, epistemic uncertain, and state.
With respect to domain type, if the user does not specify an explicit override of mixed or relaxed, Dakota infers the domain type from the method. As mentioned above, all methods currently use a mixed domain as a default, except the branch-and-bound method which is under development.

See Also

These keywords may also be of interest:

- mixed

7.4.5 continuous_design

- Keywords Area
- variables
- continuous_design

Design variable - continuous

Topics

This keyword is related to the topics:

- continuous_variables
- design_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no continuous design variables

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>Initial values for variables</td>
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<tr>
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<td>upper_bounds</td>
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<td>Specify maximum values</td>
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<td>Optional</td>
<td>scale_types</td>
<td></td>
<td>How to scale each continuous design variable</td>
</tr>
</tbody>
</table>
### Description

Continuous variables are defined by a real interval and are changed during the search for the optimal design.

#### 7.4.5.1 initial_point

- **Keywords Area**
- **variables**
- **continuous_design**
- **initial_point**

Initial values for variables

### Specification

**Alias:** cdv_initial_point  
**Argument(s):** REALLIST  
**Default:** 0.0

### Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

#### Default Behavior

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4.5.2  lower_bounds

- Keywords Area
- variables
- continuous_design
- lower_bounds

Specify minimum values

Specification

Alias: cdv_lower_bounds
Argument(s): REALLIST
Default: -infinity

Description

Specify minimum values

7.4.5.3  upper_bounds

- Keywords Area
- variables
- continuous_design
- upper_bounds

Specify maximum values

Specification

Alias: cdv_upper_bounds
Argument(s): REALLIST
Default: infinity

Description

Specify maximum values

7.4.5.4  scale_types

- Keywords Area
- variables
- continuous_design
- scale_types

How to scale each continuous design variable
7.4 variables

Specification

Alias: cdv_scale_types
Argument(s): STRINGLIST
Default: vector values = 'none'

Description

Each string in `scale_types` indicates the scaling type for each continuous design variable. They only have effect when the associated method specifies `scaling`.

The options are:

- `'value'` - characteristic value by which variables will be divided. If this is chosen, then `scales` must also be specified; 'value' is assumed if scales are given without `scale_types`
- `'auto'` - automatic scaling based on bounds.
- `'log'` - logarithmic scaling (can be used together with `scales`).

If a single string is specified it will apply to all continuous design variables. Otherwise, a scale type must be specified for each continuous design variable.

Usage Tips:

See the scaling information under specific methods, e.g., method-∗-scaling for details on how to use this keyword.

Examples

Two continuous design variables, one scaled by 4.0, the other by 0.1, then further log scaled:

```plaintext
continuous_design = 2
    initial_point -1.2  1.0
    lower_bounds  -2.0  0.001
    upper_bounds  2.0   2.0
    descriptors   'x1'  'x2'
    scale_types   'value' 'log'
    scales        4.0   0.1
```

7.4.5.5 scales

- Keywords Area
- variables
- continuous_design
- scales

Characteristic values to scale continuous design variables

Specification

Alias: cdv_scales
Argument(s): REALLIST
Default: vector values = 1. (no scaling)
Description

Each real value in scales is a nonzero characteristic value to be used in scaling each continuous design variable. They only have effect when the associated method specifies scaling.

This keyword is required for scale_types of ‘value’ and optional for ‘auto’ and ‘log’. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative design variables.

If a single real value is specified it will apply to all continuous design variables. Otherwise, a scale value must be specified for each continuous design variable.

Usage Tips:

When scales are specified, but not scale_types, the scaling type is assumed to be ‘value’ for all continuous design variables.

Use scale values of 1.0 to selectively avoid scaling a subset of the variables.

Scaling for linear constraints is applied after any continuous variable scaling.

See the scaling information under specific methods, e.g., method-∗-scaling for details on how to use this keyword.

Examples

Two continuous design variables, one scaled by the characteristic value 4.0, the other log-scaled without additional value scaling

```plaintext
continuous_design = 2
   initial_point = -1.2  1.0
   lower_bounds = -200  0.001
   upper_bounds = 200  2.0
   descriptors = 'x1'  'x2'
   scale_types = 'value'  'log'
   scales = 4.0  1.0
```

7.4.5.6 descriptors

- Keywords Area
- variables
  - continuous_design
- descriptors

Labels for the variables

Specification

Alias: cdv_descriptors

Argument(s): STRINGLIST

Default: cdv_\{i\}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.
7.4.6 discrete_design_range

- Keywords Area
- variables
- discrete_design_range

Design variable - discrete range-valued

Topics

This keyword is related to the topics:

- discrete_variables
- design_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no discrete design variables

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td></td>
<td>Initial values for variables</td>
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<td>lower_bounds</td>
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<td>Optional</td>
<td>upper_bounds</td>
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</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td></td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

These variables take on a range of integer values from the specified lower bound to the specified upper bound (integer interval). The details of how to specify this discrete variable are located on the discrete_variables page.

7.4.6.1 initial_point

- Keywords Area
- variables
- discrete_design_range
- initial_point

Initial values for variables
Specification

Alias: ddv_initial_point
Argument(s): INTEGERLIST
Default: 0

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.6.2  lower_bounds

- Keywords Area
- variables
  - discrete_design_range
  - lower_bounds

Specify minimum values

Specification

Alias: ddv_lower_bounds
Argument(s): INTEGERLIST
Default: INT_MIN

Description

Specify minimum values
7.4 variables

7.4.6.3 upper_bounds

- Keywords Area
- variables
- discrete_design_range
- upper_bounds

Specify maximum values

Specification

**Alias:** ddv_upper_bounds  
**Argument(s):** INTEGERLIST  
**Default:** INT_MAX

Description

Specify maximum values

7.4.6.4 descriptors

- Keywords Area
- variables
- discrete_design_range
- descriptors

Labels for the variables

Specification

**Alias:** ddv_descriptors  
**Argument(s):** STRINGLIST  
**Default:** ddriv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.7 discrete_design_set

- Keywords Area
- variables
- discrete_design_set

Design variable - discrete set-valued
Topics

This keyword is related to the topics:

- discrete_variables
- design_variables

Specification

Alias: none

Argument(s): none

Child Keywords:

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Dakota Keyword Description

<table>
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<tbody>
<tr>
<td>Integer-valued discrete design variables</td>
</tr>
<tr>
<td>String-valued discrete design set variables</td>
</tr>
<tr>
<td>Real-valued discrete design variables</td>
</tr>
</tbody>
</table>

Description

Discrete design variables whose values come from a set of admissible elements. Each variable specified must be of type integer, string, or real.

7.4.7.1 integer

- Keywords Area
- variables
- discrete_design_set
- integer

Integer-valued discrete design variables

Topics

This keyword is related to the topics:

- discrete_variables
- design_variables
7.4 variables

Specification

Alias: none

Argument(s): INTEGER

Default: no discrete design set integer variables

Child Keywords:

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<td>elements_per_variable</td>
<td>Number of admissible elements for each set variable</td>
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<tr>
<td>Required</td>
<td>elements</td>
<td>elements</td>
<td>The permissible values for each discrete variable</td>
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<tr>
<td>Optional</td>
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<td>Whether the set-valued variables are categorical or relaxable</td>
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<td>Optional</td>
<td>descriptors</td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

A design variable whose values come from a specified set of admissible integers. The details of how to specify this discrete variable are located on the discrete_variables page.

Examples

Four integer variables whose values will be selected from the following sets during the search for an optimal design. $y_1 \in \{0, 1\}, y_2 \in \{0, 1\}, y_3 \in \{0, 5\}$ and $y_4 \in \{10, 15, 20, 23\}$.

discrete_design_set
integer 4
descriptors 'y1' 'y2' 'y3' 'y4'
elements_per_variable 2 2 2 4
elements 0 1 0 1 0 5 10 15 20 23

7.4.7.1.1 elements_per_variable

- Keywords Area
- variables
- discrete_design_set
- integer
- elements_per_variable
Number of admissible elements for each set variable

Specification

Alias: num_set_values
Argument(s): INTEGERLIST
Default: equal distribution

Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

7.4.7.1.2 elements

• Keywords Area
• variables
• discrete_design_set
• integer
• elements

The permissible values for each discrete variable

Specification

Alias: set_values
Argument(s): INTEGERLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.7.1.3 categorical

• Keywords Area
• variables
• discrete_design_set
• integer
• categorical

Whether the set-valued variables are categorical or relaxable
7.4 variables

Specification

Alias: none
Argument(s): STRINGLIST
Child Keywords:
Description

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include ‘yes’, ‘no’, ‘true’, and ‘false’, or any abbreviation in [yYnNtTfF].

Examples

Discrete_design_set variable, ‘rotor_blades’, can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and ‘rotor_blades’ can take on any value between 2 and 7, e.g., 3, 6, or 5.5.

discrete_design_set
  integer 1
    elements 2 4 7
descriptor 'rotor_blades'
categorical 'no'

7.4.7.1.3.1 adjacency_matrix

- Keywords Area
- variables
- discrete_design_set
- integer
- categorical
- adjacency_matrix

1-0 matrix defining which categorical variable levels are related.

Specification

Alias: none

Argument(s): INTEGERLIST

Description

The adjacency_matrix keyword is associated with discrete_design_set variables that are specified to be categorical. Each such variable is associated with one $k \times k$ symmetric matrix, where $k$ is the number of values (or levels) of the variable. Entry $i, j$ of a matrix should be 1 if level $i$ and level $j$ are related by some subjective criteria or if $i = j$; it should be 0 otherwise. The matrices for all variables of the same type (string, real, or integer) are entered sequentially as a list of integers as shown in the examples below.

Default Behavior
The \texttt{adjacency_matrix} keyword is only relevant for \texttt{discrete_design_set real} and \texttt{discrete_design_set integer} variables if one or more of them have been specified to be categorical. It is always relevant for \texttt{discrete_design_set string} variables. If the user does not define an adjacency matrix, the default is method dependent. Currently, the only method that makes use of the adjacency matrix is \texttt{mesh_adaptive_search}, which uses a tri-diagonal adjacency matrix by default.

**Expected Output**

The expected output is method dependent.

**Usage Tips**

If an adjacency matrix is defined for one type of (categorical) \texttt{discrete_design_set} variable, it must be defined for all variables of that type, even for those not defined to be categorical. Those for the non-categorical set variables will be ignored.

**Examples**

The following example shows a variables specification where some real and some integer \texttt{discrete_design_set} variables are categorical.

```plaintext
variables
continuous_design = 3
  initial_point -1.0 1.5 2.0
  lower_bounds -10.0 -10.0 -10.0
  upper_bounds 10.0 10.0 10.0
  descriptors 'x1' 'x2' 'x3'
discrete_design_range = 2
  initial_point 2 2
  lower_bounds 1 1
  upper_bounds 4 9
  descriptors 'y1' 'y2'
discrete_design_set
  real = 2
    elements_per_variable = 4 5
    elements = 1.2 2.3 3.4 4.5 1.2 3.3 4.4 5.5 7.7
    descriptors 'y3' 'y4'
    categorical 'no' 'yes'
    adjacency_matrix 1 1 0 0 # Begin entry of 4x4 matrix for y3
    1 1 1 0
    0 1 1 1
    0 0 1 1
    1 0 1 0 1 # Begin entry of 5x5 matrix for y4
    1 0 1 0
    1 0 1 0
    0 1 0 1 0
    1 0 1 0

integer = 2
  elements_per_variable = 2 3
  elements = 4 7 8 9 12
  descriptors 'z1' 'z2'
  categorical 'yes' 'yes'
```

Note that for the real case, the user wants to define an adjacency matrix for the categorical variable, so adjacency matrices for both variables must be specified. The matrix for the first one will be ignored. Note that no adjacency matrix is specified for either integer categorical variable. The default will be used in both cases. Currently the only method taking advantage of adjacency matrices is \texttt{mesh_adaptive_search}, which uses a tri-diagonal adjacency matrix by default. Thus, the matrices used would be

```plaintext
z1: 1 1
   1
z2: 1 1 0
   1 1
   0 1
```

The following example shows a variables specification for string variables. Note that string variables are always
considered to be categorical. If an adjacency matrix is not specified, a method-dependent default matrix will be used.

variables,
  continuous_design = 2
  initial_point 0.5 0.5
  lower_bounds 0. 0.
  upper_bounds 1. 1.
  descriptors = 'x' 'y'
discrete_design_set string = 1
  elements = 'aniso1' 'aniso2' 'isol' 'iso2' 'iso3'
  descriptors = 'ancomp'
  adjacency_matrix
  1 1 0 0 0
  1 1 0 0 0
  0 0 1 1 1
  0 0 1 1 1
  0 0 1 1 1

See Also

These keywords may also be of interest:

- mesh_adaptive_search

7.4.7.1.4 initial_point

- Keywords Area
- variables
- discrete_design_set
- integer
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): INTEGERLIST

Default: middle set value, or rounded down

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.7.1.5 descriptors

- **Keywords Area**
- **variables**
- **discrete_design_set**
- **integer**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.7.2 string

- **Keywords Area**
- **variables**
- **discrete_design_set**
- **string**

String-valued discrete design set variables

**Topics**

This keyword is related to the topics:

- **discrete_variables**
- **design_variables**
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no discrete design set string variables

**Child Keywords:**
### Description

Discrete design variables whose values come from a specified set of admissible strings. The details of how to specify this discrete variable are located on the `discrete_variables` page. Each string element value must be quoted and may contain alphanumeric, dash, underscore, and colon. White space, quote characters, and backslash/metacharacters are not permitted.

### Examples

Two string variables whose values will be selected from the set of provided elements. The first variable, 'linear solver', takes on values from a set of three possible elements and the second variable, 'mesh_file', from a set of two possible elements.

```plaintext
discrete_design_set
string 2
  descriptors 'linear_solver' 'mesh_file'
  elements_per_variable 3 2
  elements 'cg' 'gmres' 'direct'
  'mesh64.exo' 'mesh128.exo'
```

### elements_per_variable

- **Keywords Area**
  - variables
  - discrete_design_set
  - string
  - elements_per_variable

Number of admissible elements for each set variable

---

Dakota Version 6.15 Reference Manual generated on November 5, 2021
Specification

**Alias:** num_set_values

**Argument(s):** INTEGERLIST

**Default:** equal distribution

**Description**

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

7.4.7.2.2 elements

- Keywords Area
- variables
- discrete_design_set
- string
- elements

The permissible values for each discrete variable

Specification

**Alias:** set_values

**Argument(s):** STRINGLIST

**Description**

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.7.2.3 adjacency_matrix

- Keywords Area
- variables
- discrete_design_set
- string
- adjacency_matrix

1-0 matrix defining which categorical variable levels are related.

Specification

**Alias:** none

**Argument(s):** INTEGERLIST
Description

The `adjacency_matrix` keyword is associated with `discrete_design_set` variables that are specified to be categorical. Each such variable is associated with one $k \times k$ symmetric matrix, where $k$ is the number of values (or levels) of the variable. Entry $i, j$ of a matrix should be 1 if level $i$ and level $j$ are related by some subjective criteria or if $i = j$; it should be 0 otherwise. The matrices for all variables of the same type (string, real, or integer) are entered sequentially as a list of integers as shown in the examples below.

Default Behavior

The `adjacency_matrix` keyword is only relevant for `discrete_design_set` real and `discrete_design_set` integer variables if one or more of them have been specified to be categorical. It is always relevant for `discrete_design_set` string variables. If the user does not define an adjacency matrix, the default is method dependent. Currently, the only method that makes use of the adjacency matrix is `mesh_adaptive_search`, which uses a tri-diagonal adjacency matrix by default.

Expected Output

The expected output is method dependent.

Usage Tips

If an adjacency matrix is defined for one type of (categorical) `discrete_design_set` variable, if must be defined for all variables of that type, even for those not defined to be categorical. Those for the non-categorical set variables will be ignored.

Examples

The following example shows a variables specification where some real and some integer `discrete_design_set` variables are categorical.

```plaintext
variables
continuous_design = 3
  initial_point = -1.0 1.5 2.0
  lower_bounds = -10.0 -10.0 -10.0
  upper_bounds = 10.0 10.0 10.0
descriptors = 'x1' 'x2' 'x3'
discrete_design_range = 2
  initial_point = 2 2
  lower_bounds = 1 1
  upper_bounds = 4 9
descriptors = 'y1' 'y2'
discrete_design_set
  real = 2
    elements_per_variable = 4 5
    elements = 1.2 2.3 3.4 4.5 1.2 3.3 4.4 5.5 7.7
    descriptors = 'y3' 'y4'
categorical = 'no' 'yes'
    adjacency_matrix = 1 1 0 0 # Begin entry of 4x4 matrix for y3
    1 1 1
    0 1 1
    0 0 1 1
    1 0 1 0 1 # Begin entry of 5x5 matrix for y4
    0 1 0 1 0
    1 0 1 0 1
    0 1 0 1 0
    1 0 1 0 1

integer = 2
    elements_per_variable = 2 3
    elements = 4 7 8 9 12
    descriptors = 'z1' 'z2'
categorical = 'yes' 'yes'
```

Note that for the real case, the user wants to define an adjacency matrix for the categorical variable, so adjacency matrices for both variables must be specified. The matrix for the first one will be ignored. Note that no adjacency matrix is specified for either integer categorical variable. The default will be used in both cases.
The only method taking advantage of adjacency matrices is `mesh_adaptive_search`, which uses a tri-diagonal adjacency matrix by default. Thus, the matrices used would be

\[
\begin{align*}
z_1 & : 1 & 1 \\
        & 1 & 1 \\
z_2 & : 1 & 1 & 0 \\
        & 1 & 1 & 1 \\
        & 0 & 1 & 1 \\
\end{align*}
\]

The following example shows a variables specification for string variables. Note that string variables are always considered to be categorical. If an adjacency matrix is not specified, a method-dependent default matrix will be used.

```dakota
variables,
    continuous_design = 2
    initial_point 0.5 0.5
    lower_bounds 0. 0.
    upper_bounds 1. 1.
    descriptors = 'x' 'y'
discrete_design_set string = 1
    elements = 'aniso1' 'aniso2' 'iso1' 'iso2' 'iso3'
    descriptors = 'ancomp'
    adjacency_matrix 1 1 0 0 0
                    1 1 0 0 0
                    0 0 1 1 1
                    0 0 1 1 1
                    0 0 1 1 1
```

**See Also**

These keywords may also be of interest:

- `mesh_adaptive_search`

**7.4.7.2.4 initial_point**

- **Keywords Area**
- **variables**
- **discrete_design_set**
- **string**
- **initial_point**

Initial values for variables

**Specification**

- **Alias:** none
- **Argument(s):** STRINGLIST
- **Default:** middle set value, or rounded down

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.
Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.7.2.5 descriptors

- Keywords Area
- variables
- discrete_design_set
- string
- descriptors

Labels for the variables

Specification

Alias: none

Argument(s): STRINGLIST

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.7.3 real

- Keywords Area
- variables
- discrete_design_set
- real

Real-valued discrete design variables
Topics

This keyword is related to the topics:

- discrete_variables
- design_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no discrete design set real variables

Child Keywords:

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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
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<td>elements_per_variable</td>
<td>elements_per_variable</td>
<td>Number of admissible elements for each set variable</td>
</tr>
<tr>
<td>Required</td>
<td>elements</td>
<td>elements</td>
<td>The permissible values for each discrete variable</td>
</tr>
<tr>
<td>Optional</td>
<td>categorical</td>
<td>categorical</td>
<td>Whether the set-valued variables are categorical or relaxable</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_point</td>
<td>initial_point</td>
<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

A design variable whose values come from a specified set of admissible reals. The details of how to specify this discrete variable are located on the discrete_variables page.

Examples

Two continuous, restricted variables whose values will be selected from the following sets during the search for an optimal design. \( y_1 \in \{0.25, 1.25, 2.25, 3.25, 4.25\} \), \( y_2 \in \{0, 5\} \)

```
discrete_design_set
real 2
    descriptors 'y1' 'y2'
    elements_per_variable 5 2
    elements 0.25 1.25 2.25 3.25 4.25 0 5
```
7.4.7.3.1 elements_per_variable

- Keywords Area
- variables
- discrete_design_set
- real
- elements_per_variable

Number of admissible elements for each set variable

Specification

Alias: num_set_values

Argument(s): INTEGERLIST

Default: equal distribution

Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

7.4.7.3.2 elements

- Keywords Area
- variables
- discrete_design_set
- real
- elements

The permissible values for each discrete variable

Specification

Alias: set_values

Argument(s): REALLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.
7.4.7.3.3  categorical

- Keywords Area
- variables
- discrete_design_set
- real
- categorical

Whether the set-valued variables are categorical or relaxable

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
<td></td>
<td>adjacency_matrix</td>
<td>1-0 matrix defining which categorical variable levels are related.</td>
</tr>
</tbody>
</table>

**Description**

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include 'yes', 'no', 'true', and 'false', or any abbreviation in [yYnNtTfF][.]

**Examples**

Discrete_design_set variable, 'rotor_blades', can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and 'rotor_blades' can take on any value between 2 and 7, e.g., 3, 6, or 5.5.

```plaintext
discrete_design_set
  integer 1
  elements 2 4 7
  descriptor 'rotor_blades'
  categorical 'no'
```

7.4.7.3.3.1  adjacency_matrix

- Keywords Area
- variables
- discrete_design_set
- real
- categorical
- adjacency_matrix

1-0 matrix defining which categorical variable levels are related.
Specification

**Alias:** none

**Argument(s):** INTEGERLIST

Description

The `adjacency_matrix` keyword is associated with `discrete_design_set` variables that are specified to be categorical. Each such variable is associated with one $k \times k$ symmetric matrix, where $k$ is the number of values (or levels) of the variable. Entry $i, j$ of a matrix should be 1 if level $i$ and level $j$ are related by some subjective criteria or if $i = j$; it should be 0 otherwise. The matrices for all variables of the same type (string, real, or integer) are entered sequentially as a list of integers as shown in the examples below.

Default Behavior

The `adjacency_matrix` keyword is only relevant for `discrete_design_set` real and `discrete_design_set` integer variables if one or more of them have been specified to be categorical. It is always relevant for `discrete_design_set` string variables. If the user does not define an adjacency matrix, the default is method dependent. Currently, the only method that makes use of the adjacency matrix is `mesh_adaptive_search`, which uses a tri-diagonal adjacency matrix by default.

Expected Output

The expected output is method dependent.

Usage Tips

If an adjacency matrix is defined for one type of (categorical) `discrete_design_set` variable, it must be defined for all variables of that type, even for those not defined to be categorical. Those for the non-categorical set variables will be ignored.

Examples

The following example shows a variables specification where some real and some integer `discrete_design_set` variables are categorical.

```
variables
  continuous_design = 3
    initial_point -1.0 1.5 2.0
    lower_bounds -10.0 -10.0 -10.0
    upper_bounds 10.0 10.0 10.0
    descriptors 'x1' 'x2' 'x3'
  discrete_design_range = 2
    initial_point 2 2
    lower_bounds 1 1
    upper_bounds 4 9
    descriptors 'y1' 'y2'
  discrete_design_set
    real = 2
      elements_per_variable = 4 5
      elements = 1.2 2.3 3.4 4.5 1.2 3.3 4.4 5.5 7.7
      descriptors 'y3' 'y4'
      categorical 'no' 'yes'
      adjacency_matrix 1 1 0 0 # Begin entry of 4x4 matrix for y3
        1 1 1 1
        0 1 0 0
        1 0 1 0
      # Begin entry of 5x5 matrix for y4
        1 0 1 0
        1 0 1 0
        0 1 0 1
        1 0 1 0
      integer = 2
        elements_per_variable = 2 3
        elements = 4 7 8 9 12
```
Note that for the real case, the user wants to define an adjacency matrix for the categorical variable, so adjacency matrices for both variables must be specified. The matrix for the first one will be ignored. Note that no adjacency matrix is specified for either integer categorical variable. The default will be used in both cases. Currently the only method taking advantage of adjacency matrices is `mesh_adaptive_search`, which uses a tri-diagonal adjacency matrix by default. Thus, the matrices used would be

\[
\begin{bmatrix}
1 & 1 \\
1 & 1 \\
\end{bmatrix}
\]
\[
\begin{bmatrix}
1 & 1 & 0 \\
1 & 1 & 1 \\
0 & 1 & 1 \\
\end{bmatrix}
\]

The following example shows a variables specification for string variables. Note that string variables are always considered to be categorical. If an adjacency matrix is not specified, a method-dependent default matrix will be used.

```plaintext
variables,
  continuous_design = 2
  initial_point 0.5 0.5
  lower_bounds 0. 0.
  upper_bounds 1. 1.
  descriptors = 'x' 'y'
  discrete_design_set string = 1
  elements = 'aniso1' 'aniso2' 'iso1' 'iso2' 'iso3'
  descriptors = 'ancomp'
  adjacency_matrix 1 1 0 0 0
                  1 1 0 0 0
                  0 0 1 1 1
                  0 0 1 1 1
                  0 0 1 1 1
```

See Also

These keywords may also be of interest:

- `mesh_adaptive_search`

7.4.7.3.4 initial_point

- `Keywords Area`
- `variables`
- `discrete_design_set`
- `real`
- `initial_point`

Initial values for variables

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** middle set value, or rounded down
Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.7.3.5 descriptors

- Keywords Area
- variables
- discrete_design_set
- real
- descriptors

Labels for the variables

Specification

Alias: none

Argument(s): STRINGLIST

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.8 normal_uncertain

- Keywords Area
- variables
- normal_uncertain

Aleatory uncertain variable - normal (Gaussian)
This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Alias: none

Default: no normal uncertain variables

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
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<tr>
<td>Required</td>
<td>means</td>
<td>means</td>
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</tr>
<tr>
<td>Required</td>
<td>std_deviations</td>
<td>std_deviations</td>
<td>Second parameter of the distribution</td>
</tr>
<tr>
<td>Optional</td>
<td>lower_bounds</td>
<td>lower_bounds</td>
<td>Specify minimum values</td>
</tr>
<tr>
<td>Optional</td>
<td>upper_bounds</td>
<td>upper_bounds</td>
<td>Specify maximum values</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_point</td>
<td>initial_point</td>
<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

The number of normal uncertain variables, their means, and standard deviations are required specifications, while the distribution lower and upper bounds and variable descriptors are optional specifications. The normal distribution is widely used to model uncertain variables such as population characteristics. It is also used to model the mean of a sample: as the sample size becomes very large, the Central Limit Theorem states that the distribution of the mean becomes approximately normal, regardless of the distribution of the original variables.

The density function for the normal distribution is:

\[
f(x) = \frac{1}{\sqrt{2\pi} \sigma} \exp \left( -\frac{1}{2} \left( \frac{x - \mu}{\sigma} \right)^2 \right)
\]

where \( \mu \) and \( \sigma \) are the mean and standard deviation of the normal distribution, respectively.

Note that if you specify bounds for a normal distribution, the sampling occurs from the underlying distribution with the given mean and standard deviation, but samples are not taken outside the bounds (see "bounded normal" distribution type in[93]). This can result in the mean and the standard deviation of the sample data being different from the mean and standard deviation of the underlying distribution. For example, if you are sampling from a normal distribution with a mean of 5 and a standard deviation of 3, but you specify bounds of 1 and 7, the resulting mean of the samples will be around 4.3 and the resulting standard deviation will be around 1.6. This is because you have bounded the original distribution significantly, and asymmetrically, since 7 is closer to the original mean than 1.
7.4 variables

Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be $[\mu - 3\sigma, \mu + 3\sigma]$.

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means, correcting to bounds if needed.

7.4.8.1 means

- Keywords Area
- variables
- normal_uncertain
- means

First parameter of the distribution

Specification

Alias: nuv_means
Argument(s): REALLIST

Description

Means

7.4.8.2 std_deviations

- Keywords Area
- variables
- normal_uncertain
- std_deviations

Second parameter of the distribution

Specification

Alias: nuv_std_deviations
Argument(s): REALLIST

Description

Standard deviation
7.4.8.3 lower_bounds

• Keywords Area
• variables
• normal_uncertain
• lower_bounds

Specify minimum values

Specification

Alias: nuv_lower_bounds

Argument(s): REALLIST

Default: -infinity

Description

Specify minimum values

7.4.8.4 upper_bounds

• Keywords Area
• variables
• normal_uncertain
• upper_bounds

Specify maximum values

Specification

Alias: nuv_upper_bounds

Argument(s): REALLIST

Default: infinity

Description

Specify maximum values

7.4.8.5 initial_point

• Keywords Area
• variables
• normal_uncertain
• initial_point

Initial values for variables
7.4 variables

Specification

**Alias:** none

**Argument(s):** REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.8.6 descriptors

- Keywords Area
- variables
- normal_uncertain
- descriptors

Labels for the variables

Specification

**Alias:** nuv_descriptors

**Argument(s):** STRINGLIST

**Default:** nuv_[i]

Description

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.
7.4.9  lognormal_uncertain

- Keywords Area
- variables
- lognormal_uncertain

Aleatory uncertain variable - lognormal

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no lognormal uncertain variables

**Child Keywords:**

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<td>lambdas</td>
<td>First parameter of the lognormal distribution (option 3)</td>
</tr>
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<td>First parameter of the lognormal distribution (options 1 &amp; 2)</td>
</tr>
<tr>
<td>Optional</td>
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<td>Specify minimum values</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>upper_bounds</td>
<td>Specify maximum values</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>initial_point</td>
<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

If the logarithm of an uncertain variable $X$ has a normal distribution, that is $\log X \sim N(\mu, \sigma)$, then $X$ is distributed with a lognormal distribution. The lognormal is often used to model:

1. time to perform some task
2. variables which are the product of a large number of other quantities, by the Central Limit Theorem
3. quantities which cannot have negative values.
The number of lognormal uncertain variables, their means, and either standard deviations or error factors must be specified, while the distribution lower and upper bounds and variable descriptors are optional specifications. These distribution bounds can be used to truncate the tails of lognormal distributions, which as for bounded normal, can result in the mean and the standard deviation of the sample data being different from the mean and standard deviation of the underlying distribution (see "bounded lognormal" and "bounded lognormal-n" distribution types in [93]).

For the lognormal variables, one may specify either the mean $\mu$ and standard deviation $\sigma$ of the actual lognormal distribution (option 1), the mean $\mu$ and error factor $\varepsilon$ of the actual lognormal distribution (option 2), or the mean $\lambda$ ("lambda") and standard deviation $\zeta$ ("zeta") of the underlying normal distribution (option 3).

The conversion equations from lognormal mean $\mu$ and either lognormal error factor $\varepsilon$ or lognormal standard deviation $\sigma$ to the mean $\lambda$ and standard deviation $\zeta$ of the underlying normal distribution are as follows:

$$
\zeta = \frac{\ln(\varepsilon)}{1.645}
$$

$$
\zeta^2 = \ln \left( \frac{\sigma^2}{\mu^2} + 1 \right)
$$

$$
\lambda = \ln(\mu) - \frac{\zeta^2}{2}
$$

Conversions from $\lambda$ and $\zeta$ back to $\mu$ and $\varepsilon$ or $\sigma$ are as follows:

$$
\mu = \exp \left( \lambda + \frac{\zeta^2}{2} \right)
$$

$$
\sigma^2 = \exp \left( 2\lambda + \zeta^2 \right) \left( \exp \left( \zeta^2 \right) - 1 \right)
$$

$$
\varepsilon = \exp (1.645\zeta)
$$

The density function for the lognormal distribution is:

$$
f(x) = \frac{1}{\sqrt{2\pi}\zeta x} \exp \left( -\frac{1}{2} \left( \frac{\ln(x) - \lambda}{\zeta} \right)^2 \right)
$$

**Theory**

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be $[0, \mu + 3\sigma]$.

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means, correcting to bounds if needed.

**7.4.9.1 lambdas**

- **Keywords Area**
- **variables**
- **lognormal_uncertain**
- **lambdas**

First parameter of the lognormal distribution (option 3)
Specification

**Alias:** lnuv_lambdas

**Argument(s):** REALLIST

**Child Keywords:**
7.4 variables

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required</td>
<td></td>
<td>zetas</td>
<td>Second parameter of the lognormal distribution (option 3)</td>
</tr>
</tbody>
</table>

Description

For the lognormal variables, one may specify the mean \( \lambda \) ("lambda") and standard deviation \( \zeta \) ("zeta") of the underlying normal distribution.

7.4.9.1.1 zetas

- Keywords Area
- variables
- lognormal_uncertain
- lambdas
- zetas

Second parameter of the lognormal distribution (option 3)

Specification

**Alias:** lnv_zetas

**Argument(s):** REALLIST

Description

For the lognormal variables, one may specify the mean \( \lambda \) ("lambda") and standard deviation \( \zeta \) ("zeta") of the underlying normal distribution.

7.4.9.2 means

- Keywords Area
- variables
- lognormal_uncertain
- means

First parameter of the lognormal distribution (options 1 & 2)

Specification

**Alias:** lnv_means

**Argument(s):** REALLIST

**Child Keywords:**
For the lognormal variables, one may specify either the mean $\mu$ and standard deviation $\sigma$ of the actual lognormal distribution, the mean $\mu$ and error factor $\varepsilon$ of the actual lognormal distribution.

This corresponds to the mean of the lognormal random variable.

**7.4.9.2.1 std_deviations**

- **Keywords Area**
- **variables**
- **lognormal_uncertain**
- **means**
- **std_deviations**

Second parameter of the lognormal distribution (option 1)

**Specification**

**Alias:** lnuv_std_deviations

**Argument(s):** REALLIST

**Description**

For the lognormal variables, one may specify either the mean $\mu$ and standard deviation $\sigma$ of the actual lognormal distribution.

This corresponds to the standard deviation of the lognormal random variable.

**7.4.9.2.2 error_factors**

- **Keywords Area**
- **variables**
- **lognormal_uncertain**
- **means**
- **error_factors**

Second parameter of the lognormal distribution (option 2)
7.4 variables

Specification

**Alias:** lnuv_error_factors

**Argument(s):** REALLIST

Description

For the lognormal variables, one may specify the mean $\mu$ and error factor $\varepsilon$ of the actual lognormal distribution. This specifies the error function of the lognormal random variable.

7.4.9.3 lower_bounds

- **Keywords** Area
- **variables**
- **lognormal_uncertain**
- **lower_bounds**

Specify minimum values

Specification

**Alias:** lnuv_lower_bounds

**Argument(s):** REALLIST

**Default:** 0

Description

Specify minimum values

7.4.9.4 upper_bounds

- **Keywords** Area
- **variables**
- **lognormal_uncertain**
- **upper_bounds**

Specify maximum values

Specification

**Alias:** lnuv_upper_bounds

**Argument(s):** REALLIST

**Default:** infinity
Description

Specify maximum values

7.4.9.5  initial_point

- Keywords Area
- variables
- lognormal_uncertain
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.9.6  descriptors

- Keywords Area
- variables
- lognormal_uncertain
- descriptors

Labels for the variables
7.4 variables

Specification

Alias: lnuv_descriptors

Argument(s): STRINGLIST

Default: lnuv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.10 uniform_uncertain

- Keywords Area
- variables
- uniform_uncertain

Aleatory uncertain variable - uniform

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no uniform uncertain variables

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Required</td>
<td>lower_bounds</td>
<td></td>
<td>Specify minimum values</td>
</tr>
<tr>
<td>Required</td>
<td>upper_bounds</td>
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<tr>
<td>Optional</td>
<td>initial_point</td>
<td></td>
<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td></td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>
The number of uniform uncertain variables and their distribution lower and upper bounds are required specifications, while variable descriptors is an optional specification. The uniform distribution has the density function:

\[ f(x) = \frac{1}{U-L} \]

where \( U \) and \( L \) are the upper and lower bounds of the uniform distribution, respectively. The mean of the uniform distribution is \( \frac{U+L}{2} \) and the variance is \( \frac{(U-L)^2}{12} \).

Theory

This distribution is a special case of the more general beta distribution.

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.10.1 lower_bounds

- Keywords Area
- variables
- uniform_uncertain
- lower_bounds

Specify minimum values

Specification

Alias: uuv_lower_bounds

Argument(s): REALLIST

Description

Specify minimum values

7.4.10.2 upper_bounds

- Keywords Area
- variables
- uniform_uncertain
- upper_bounds

Specify maximum values

Specification

Alias: uuv_upper_bounds

Argument(s): REALLIST
Description

Specify maximum values

7.4.10.3 initial_point

- **Keywords Area**
- **variables**
- **uniform_uncertain**
- **initial_point**

Initial values for variables

Specification

**Alias:** none

**Argument(s):** REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including *continuous_design*, *continuous_state*, *discrete_design_range*, and *discrete_state_range* are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including *discrete_design_set* and *discrete_state_set* default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.10.4 descriptors

- **Keywords Area**
- **variables**
- **uniform_uncertain**
- **descriptors**

Labels for the variables
Specification

Alias: uuv_descriptors

Argument(s): STRINGLIST

Default: uuv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.11 loguniform_uncertain

- Keywords Area
- variables
- loguniform_uncertain

Aleatory uncertain variable - loguniform

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no loguniform uncertain variables

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td></td>
<td>Labels for the variables</td>
</tr>
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</table>
Description

If the logarithm of an uncertain variable $X$ has a uniform distribution, that is $\log X \sim U(L_{LU}, U_{LU})$, then $X$ is distributed with a loguniform distribution. The distribution lower bound is $L_{LU}$ and upper bound is $U_{LU}$. The loguniform distribution has the density function:

$$f(x) = \frac{1}{x(\ln(U_{LU}) - \ln(L_{LU}))}$$

Theory

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.11.1 lower_bounds

- Keywords Area
- variables
- loguniform_uncertain
- lower_bounds

Specify minimum values

 Specification

Alias: luuv_lower_bounds

Argument(s): REALLIST

Description

Specify minimum values

7.4.11.2 upper_bounds

- Keywords Area
- variables
- loguniform_uncertain
- upper_bounds

Specify maximum values

 Specification

Alias: luuv_upper_bounds

Argument(s): REALLIST

Description

Specify maximum values
7.4.11.3 initial_point

- Keywords Area
- variables
  - loguniform_uncertain
  - initial_point

Initial values for variables

Specification

Alias: none
Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.11.4 descriptors

- Keywords Area
- variables
  - loguniform_uncertain
  - descriptors

Labels for the variables

Specification

Alias: luuv_descriptors
Argument(s): STRINGLIST
Default: luuv_{i}
Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.12 triangular_uncertain

- Keywords Area
- variables
- triangular_uncertain

Aleatory uncertain variable - triangular

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no triangular uncertain variables

Child Keywords:

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</tr>
<tr>
<td></td>
<td>descriptors</td>
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<td>variables</td>
</tr>
</tbody>
</table>

Description

The triangular distribution is often used when one does not have much data or information, but does have an estimate of the most likely value and the lower and upper bounds. The number of triangular uncertain variables, the modes, and the distribution lower and upper bounds are required specifications, while variable descriptors is an optional specification.
The density function for the triangular distribution is:

\[ f(x) = \frac{2(x - L)}{(U - L)(M - L)} \]

if \( L \leq x \leq M \), and

\[ f(x) = \frac{2(U - x)}{(U - L)(U - M)} \]

if \( M \leq x \leq U \), and 0 elsewhere. In these equations, \( L \) is the lower bound, \( U \) is the upper bound, and \( M \) is the mode of the triangular distribution.

7.4.12.1 modes

- Keywords Area
- variables
- triangular_uncertain
- modes

Distribution parameter

Specification

Alias: tuv_modes
Argument(s): REALLIST

Description

Specify the modes

7.4.12.2 lower_bounds

- Keywords Area
- variables
- triangular_uncertain
- lower_bounds

Specify minimum values

Specification

Alias: tuv_lower_bounds
Argument(s): REALLIST

Description

Specify minimum values
7.4 variables

7.4.12.3 upper_bounds

• Keywords Area
• variables
• triangular_uncertain
• upper_bounds

Specify maximum values

Specification

Alias: tuv_upper_bounds
Argument(s): REALLIST

Description

Specify maximum values

7.4.12.4 initial_point

• Keywords Area
• variables
• triangular_uncertain
• initial_point

Initial values for variables

Specification

Alias: none
Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

• Mean/midpoint (or next smaller integer value) when doubly-bounded

• 0 when unbounded

• 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.12.5 descriptors

- **Keywords Area**
- **variables**
- **triangular_uncertain**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** `tuv_descriptors`

**Argument(s):** STRINGLIST

**Default:** `tuv_{i}`

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

### 7.4.13 exponential_uncertain

- **Keywords Area**
- **variables**
- **exponential_uncertain**

Aleatory uncertain variable - exponential

**Topics**

This keyword is related to the topics:

- `continuous_variables`
- `aleatory_uncertain_variables`
Specification

**Alias**: none

**Argument(s)**: INTEGER

**Default**: no exponential uncertain variables

**Child Keywords**: 
### Description

The exponential distribution is often used for modeling failure rates.

The density function for the exponential distribution is given by:

\[ f(x) = \frac{1}{\beta} \exp \left( -\frac{x}{\beta} \right) \]

where \( \mu = \beta \) and \( \sigma^2 = \beta^2 \).

Note that this distribution is a special case of the more general gamma distribution.

### Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be \([0, \mu + 3\sigma]\).

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.13.1 betas

- Keywords Area
- variables
- exponential_uncertain
- betas

Parameter of the exponential distribution

### Specification

**Alias:** euv_betas

**Argument(s):** REALLIST

**Description**

Specifies the list of \( \beta \) parameters to define the distributions of the exponential random variables. Length must match the other parameters and the number of exponential random variables.

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</table>
7.4 variables

7.4.13.2 initial_point

- Keywords Area
- variables
- exponential_uncertain
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.13.3 descriptors

- Keywords Area
- variables
- exponential_uncertain
- descriptors

Labels for the variables

Specification

Alias: euv_descriptors

Argument(s): STRINGLIST

Default: euv_{i}
Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.14  beta_uncertain

- Keywords Area
- variables
- beta_uncertain

Aleatory uncertain variable - beta

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no beta uncertain variables

**Child Keywords:**

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</tr>
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<td>Optional</td>
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<td></td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

The number of beta uncertain variables, the alpha and beta parameters, and the distribution upper and lower bounds are required specifications, while the variable descriptors is an optional specification. The beta distribution can be
helpful when the actual distribution of an uncertain variable is unknown, but the user has a good idea of the bounds, the mean, and the standard deviation of the uncertain variable. The density function for the beta distribution is

\[
f(x) = \frac{\Gamma(\alpha + \beta)}{\Gamma(\alpha)\Gamma(\beta)} \frac{(x-L)^{\alpha-1}(U-x)^{\beta-1}}{(U-L)^{\alpha+\beta-1}}
\]

where \( \Gamma(\alpha) \) is the gamma function and \( B(\alpha, \beta) = \frac{\Gamma(\alpha)\Gamma(\beta)}{\Gamma(\alpha + \beta)} \) is the beta function. To calculate the mean and standard deviation from the alpha, beta, upper bound, and lower bound parameters of the beta distribution, the following expressions may be used.

\[
\mu = L + \frac{\alpha}{\alpha + \beta}(U - L)
\]

\[
\sigma^2 = \frac{\alpha \beta}{(\alpha + \beta)^2(\alpha + \beta + 1)}(U - L)^2
\]

Solving these for \( \alpha \) and \( \beta \) gives:

\[
\alpha = (\mu - L) \frac{(\mu - L)(U - \mu) - \sigma^2}{\sigma^2(U - L)}
\]

\[
\beta = (U - \mu) \frac{(\mu - L)(U - \mu) - \sigma^2}{\sigma^2(U - L)}
\]

Note that the uniform distribution is a special case of this distribution for parameters \( \alpha = \beta = 1 \).

**Theory**

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

### 7.4.14.1 alphas

- **Keywords Area**
- **variables**
- **beta_uncertain**
- **alphas**

First parameter of the beta distribution

**Specification**

**Alias:** buv_alphas

**Argument(s):** REALLIST

**Description**

Specifies the list of \( \alpha \) parameters to define the distributions of the beta random variables. Length must match the other parameters and the number of beta random variables.
7.4.14.2 betas

- Keywords Area
- variables
- beta_uncertain
- betas

Second parameter of the beta distribution

**Specification**

**Alias:** buv_betas

**Argument(s):** REALLIST

**Description**

Specifies the list of \( \beta \) parameters to define the distributions of the beta random variables. Length must match the other parameters and the number of beta random variables.

7.4.14.3 lower_bounds

- Keywords Area
- variables
- beta_uncertain
- lower_bound

Specify minimum values

**Specification**

**Alias:** buv_lower_bounds

**Argument(s):** REALLIST

**Description**

Specify minimum values

7.4.14.4 upper_bounds

- Keywords Area
- variables
- beta_uncertain
- upper_bound

Specify maximum values
7.4 variables

Specification

**Alias:** buv_upper_bounds

**Argument(s):** REALLIST

Description

Specify maximum values

7.4.14.5 initial_point

- **Keywords Area**
- **variables**
- **beta_uncertain**
- **initial_point**

Initial values for variables

Specification

**Alias:** none

**Argument(s):** REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.14.6 descriptors

- **Keywords Area**
- **variables**
• beta_uncertain
• descriptors

Labels for the variables

Specification

Alias: buv_descriptors
Argument(s): STRINGLIST
Default: buv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.15 gamma_uncertain

• Keywords Area
• variables
• gamma_uncertain

Aleatory uncertain variable - gamma

Topics

This keyword is related to the topics:

• continuous_variables
• aleatory_uncertain_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no gamma uncertain variables

Child Keywords:

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</table>
Description

The gamma distribution is sometimes used to model time to complete a task, such as a repair or service task. It is a very flexible distribution with its shape governed by alpha and beta.

The density function for the gamma distribution is given by:

\[ f(x) = \frac{x^{\alpha-1} \exp \left( \frac{-x}{\beta} \right)}{\beta^\alpha \Gamma(\alpha)} . \]

\[ \mu = \alpha \beta, \quad \text{and} \quad \sigma^2 = \alpha \beta^2. \]
Note that the exponential distribution is a special case of this distribution for parameter \( \alpha = 1 \).

Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be \([0, \mu + 3\sigma]\).

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.15.1 alphas

- **Keywords** Area
- **variables**
- **gamma_uncertain**
- **alphas**

First parameter of the gamma distribution

Specification

**Alias:** gauv_alphas

**Argument(s):** REALLIST

**Description**

Specifies the list of \( \alpha \) parameters to define the distributions of the gamma random variables. Length must match the other parameters and the number of gamma random variables.
7.4.15.2 betas

- Keywords Area
- variables
- gamma_uncertain
- betas

Second parameter of the gamma distribution

Specification

**Alias:** gauv_betas  
**Argument(s):** REALLIST

Description

Specifies the list of $\beta$ parameters to define the distributions of the gamma random variables. Length must match the other parameters and the number of gamma random variables.

7.4.15.3 initial_point

- Keywords Area
- variables
- gamma_uncertain
- initial_point

Initial values for variables

Specification

**Alias:** none  
**Argument(s):** REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.15.4 descriptors

- **Keywords Area**
- **variables**
- **gamma_uncertain**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** `gauv_descriptors`

**Argument(s):** STRINGLIST

**Default:** `gauv_{i}`

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

### 7.4.16 gumbel_uncertain

- **Keywords Area**
- **variables**
- **gumbel_uncertain**

Aleatory uncertain variable - gumbel

**Topics**

This keyword is related to the topics:

- **continuous_variables**
- **aleatory_uncertain_variables**
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no gumbel uncertain variables

**Child Keywords:**
7.4 variables

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<td>Labels for the variables</td>
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Description

The Gumbel distribution is also referred to as the Type I Largest Extreme Value distribution. The distribution of maxima in sample sets from a population with a normal distribution will asymptotically converge to this distribution. It is commonly used to model demand variables such as wind loads and flood levels.

The density function for the Gumbel distribution is given by:

\[
f(x) = \alpha \exp(-\alpha(x - \beta)) \exp\left(-e^{-\alpha(x - \beta)}\right),
\]

\[
\mu = \beta + \frac{0.5772}{\alpha}, \quad \text{and} \quad \sigma = \frac{\pi}{\sqrt{6} \alpha}.
\]

Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be \([\mu - 3\sigma, \mu + 3\sigma]\).

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.16.1 alphas

- **Keywords Area**
- **variables**
- **gumbel_uncertain**
- **alphas**

First parameter of the gumbel distribution

Specification

**Alias:** guuv_alphas

**Argument(s):** REALLIST

Description

Specifies the list of \(\beta\) parameters to define the distributions of the gumbel random variables. Length must match the other parameters and the number of gumbel random variables.
7.4.16.2 betas

- Keywords Area
- variables
- gumbel_uncertain
- betas

Second parameter of the gumbel distribution

Specification

**Alias**: guuv_betas

**Argument(s)**: REALLIST

**Description**

Specifies the list of \( \beta \) parameters to define the distributions of the gumbel random variables. Length must match the other parameters and the number of gumbel random variables.

7.4.16.3 initial_point

- Keywords Area
- variables
- gumbel_uncertain
- initial_point

Initial values for variables

Specification

**Alias**: none

**Argument(s)**: REALLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### descriptors

- **Keywords Area**
- **variables**
- **gumbel_uncertain**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** `guuv_descriptors`

**Argument(s):** STRINGLIST

**Default:** `guuv_{i}`

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

### frechet_uncertain

- **Keywords Area**
- **variables**
- **frechet_uncertain**

Aleatory uncertain variable - Frechet

**Topics**

This keyword is related to the topics:

- **continuous_variables**
- **aleatory_uncertain_variables**
Specification

Alias: none
Argument(s): INTEGER
Default: no frechet uncertain variables
Child Keywords:
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Description

The Frechet distribution is also referred to as the Type II Largest Extreme Value distribution. The distribution of maxima in sample sets from a population with a lognormal distribution will asymptotically converge to this distribution. It is commonly used to model non-negative demand variables.

The density function for the frechet distribution is:

\[ f(x) = \frac{\alpha}{\beta} \left( \frac{\beta}{x} \right)^{\alpha + 1} \exp \left( - \left( \frac{\beta}{x} \right)^\alpha \right), \]

\[ \mu = \beta \Gamma(1 - \frac{1}{\alpha}), \text{ and } \sigma^2 = \beta^2 \left[ \Gamma(1 - \frac{2}{\alpha}) - \Gamma^2(1 - \frac{1}{\alpha}) \right] \]

Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be \([0, \mu + 3\sigma]\).

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.17.1 alphas

- Keywords Area
- variables
- frechet_uncertain
- alphas

First parameter of the Frechet distribution

Specification

**Alias:** fuv_alphas

**Argument(s):** REALLIST

Description

Specifies the list of \(\alpha\) parameters to define the distributions of the Frechet random variables. Length must match the other parameters and the number of Frechet random variables.
7.4.17.2  betas

- Keywords Area
- variables
- frechet_uncertain
- betas

Second parameter of the Frechet distribution

**Specification**

**Alias:** fuv_betas  
**Argument(s):** REALLIST

**Description**

Specifies the list of $\beta$ parameters to define the distributions of the Frechet random variables. Length must match the other parameters and the number of Frechet random variables.

7.4.17.3  initial_point

- Keywords Area
- variables
- frechet_uncertain
- initial_point

Initial values for variables

**Specification**

**Alias:** none  
**Argument(s):** REALLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including \texttt{discrete\_design\_set} and \texttt{discrete\_state\_set} default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.17 descriptors

- Keywords Area
- variables
- frechet\_uncertain
- descriptors

Labels for the variables

Specification

\textbf{Alias:} fuv\_descriptors

\textbf{Argument(s):} STRINGLIST

\textbf{Default:} fuv\_{\{i\}}

Description

The optional variable labels specification \texttt{descriptors} is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.18 weibull\_uncertain

- Keywords Area
- variables
- weibull\_uncertain

Aleatory uncertain variable - Weibull

Topics

This keyword is related to the topics:

- \texttt{continuous\_variables}
- \texttt{aleatory\_uncertain\_variables}
Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no weibull uncertain variables

**Child Keywords:**
Description

The Weibull distribution is also referred to as the Type III Smallest Extreme Value distribution. The Weibull distribution is commonly used in reliability studies to predict the lifetime of a device. It is also used to model capacity variables such as material strength.

The density function for the Weibull distribution is given by:

\[ f(x) = \frac{\alpha}{\beta} \left(\frac{x}{\beta}\right)^{\alpha-1} \exp\left(-\left(\frac{x}{\beta}\right)^\alpha\right), \]

\[ \mu = \beta \Gamma\left(1 + \frac{1}{\alpha}\right), \text{ and } \sigma = \mu \sqrt{\frac{\Gamma(1+\frac{2}{\alpha})}{\Gamma^2(1+\frac{1}{\alpha})} - 1} \]

7.4.18.1 alphas

- Keywords Area
- variables
- weibull_uncertain
- alphas

First parameter of the Weibull distribution

Specification

Alias: wuv_alphas

Argument(s): REALLIST

Description

Specifies the list of \( \alpha \) parameters to define the distributions of the Weibull random variables. Length must match the other parameters and the number of Weibull random variables.

7.4.18.2 betas

- Keywords Area
Second parameter of the Weibull distribution

**Specification**

**Alias:** wuv_betas  
**Argument(s):** REALLIST

**Description**

Specifies the list of $\beta$ parameters to define the distributions of the Weibull random variables. Length must match the other parameters and the number of Weibull random variables.

7.4.18.3  initial_point

- **Keywords Area**
- **variables**
- **weibull_uncertain**
- **initial_point**

Initial values for variables

**Specification**

**Alias:** none  
**Argument(s):** REALLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4 variables

7.4.18.4 descriptors

- Keywords Area
- variables
- weibull_uncertain
- descriptors

Labels for the variables

Specification

Alias: wuv_descriptors
Argument(s): STRINGLIST
Default: wuv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.19 histogram_bin_uncertain

- Keywords Area
- variables
- histogram_bin_uncertain

Aleatory uncertain variable - continuous histogram

Topics

This keyword is related to the topics:

- continuous_variables
- aleatory_uncertain_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no histogram bin uncertain variables

Child Keywords:
### Dakota Keyword Description

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>Real abscissas for a bin histogram</td>
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<td>initial_point</td>
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</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td></td>
<td>Labels for the variables</td>
</tr>
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</table>

### Description

Histogram uncertain variables are typically used to model a set of empirical data. The bin histogram (contrast: histogram_point_uncertain) is a continuous aleatory distribution characterized by bins of non-zero width where the uncertain variable may lie, together with the relative frequencies of each bin. Hence it can be used to specify a marginal probability density function arising from data.

The histogram_bin_uncertain keyword specifies the number of variables to be characterized as continuous histograms. The required sub-keywords are: abscessas (ranges of values the variable can take on) and either ordinates or counts (characterizing each variable’s frequency information). When using histogram bin variables, each variable must be defined by at least one bin (with two bounding value pairs). When more than one histogram bin variable is active, pairs_per_variable can be used to specify unequal apportionment of provided bin pairs among the variables.

The abscessas specification defines abscissa values ("x" coordinates) for the probability density function of each histogram variable. When paired with counts, the specifications provide sets of (x,c) pairs for each histogram variable where c defines a count (i.e., a frequency or relative probability) associated with a bin. If using bins of unequal width and specification of probability densities is more natural, then the counts specification can be replaced with an ordinates specification ("y" coordinates) in order to support interpretation of the input as (x,y) pairs defining the profile of a "skyline" probability density function.

Conversion between the two specifications is straightforward: a count/frequency is a cumulative probability quantity defined from the product of the ordinate density value and the x bin width. Thus, in the cases of bins of equal width, ordinate and count specifications are equivalent. In addition, ordinates and counts may be relative values; it is not necessary to scale them as all user inputs will be normalized.

To fully specify a bin-based histogram with n bins (potentially of unequal width), n+1 (x,c) or (x,y) pairs must be specified with the following features:

- x is the parameter value for the left boundary of a histogram bin and c is the corresponding count for that bin. Alternatively, y defines the ordinate density value for this bin within a skyline probability density function. The right boundary of the bin is defined by the left boundary of the next pair.

- the final pair specifies the right end of the last bin and must have a c or y value of zero.
• the \( x \) values must be strictly increasing.

• all \( c \) or \( y \) values must be positive, except for the last which must be zero.

• a minimum of two pairs must be specified for each bin-based histogram variable.

Examples

The \texttt{pairs_per_variable} specification provides for the proper association of multiple sets of \((x,c)\) or \((x,y)\) pairs with individual histogram variables. For example, in this input snippet

\begin{verbatim}
histogram_bin_uncertain = 2
pairs_per_variable = 3  4
abscissas = 5  8  10 .1 .2 .3 .4
counts = 17  21  0 12 24 12  0
descriptors = 'hbu_1' 'hbu_2'
\end{verbatim}

\texttt{pairs_per_variable} associates the first 3 \((x,c)\) pairs from \texttt{abscissas} and \texttt{counts} \{(5,17),(8,21),(10,0)\} with one bin-based histogram variable, where one bin is defined between 5 and 8 with a count of 17 and another bin is defined between 8 and 10 with a count of 21. The following set of 4 \((x,c)\) pairs \{(.1,12),(.2,24),(.3,12),(.4,0)\} defines a second bin-based histogram variable containing three equal-width bins with counts 12, 24, and 12 (middle bin is twice as probable as the other two).

See Also

These keywords may also be of interest:

• \texttt{histogram_point_uncertain}

FAQ

\textbf{Difference between bin and point histograms:} A (continuous) bin histogram specifies bins of non-zero width, whereas a (discrete) point histogram specifies individual point values, which can be thought of as bins with zero width. In the terminology of LHS\cite{LHS93}, the bin pairs specification defines a "continuous linear" distribution and the point pairs specification defines a "discrete histogram" distribution (although the points are real-valued, the number of possible values is finite).

7.4.19.1 \texttt{pairs_per_variable}

• \texttt{Keywords Area}

• \texttt{variables}

• \texttt{histogram_bin_uncertain}

• \texttt{pairs_per_variable}

Number of pairs defining each histogram bin variable

Specification

\textbf{Alias:} \texttt{num_pairs}

\textbf{Argument(s):} \texttt{INTEGERLIST}

\textbf{Default:} equal distribution
Description

By default, the list of abscissas and counts or ordinates will be evenly divided among the histogram_bin_uncertain variables. pairs_per_variable is a list of integers that specify the number of pairs to apportion to each variable.

7.4.19.2 abscissas

- Keywords Area
- variables
- histogram_bin_uncertain
- abscissas

Real abscissas for a bin histogram

Specification

Alias: huv_bin_abscissas
Argument(s): REALLIST

Description

A list of real abscissa ("x" coordinate) values characterizing the probability density function for each of the histogram_bin_uncertain variables. These are paired with either counts or ordinates. See histogram_bin_uncertain for details and examples.

7.4.19.3 ordinates

- Keywords Area
- variables
- histogram_bin_uncertain
- ordinates

Ordinates specifying a "skyline" probability density function

Specification

Alias: huv_bin_ordinates
Argument(s): REALLIST

Description

The ordinates list of real values defines the profile of a "skyline" probability density function by pairing with the specified abscissas. See histogram_bin_uncertain for details.
7.4 variables

7.4.19.4 counts

- Keywords Area
- variables
- histogram_bin_uncertain
- counts

Frequency or relative probability of each bin

Specification

Alias: huv_bin_counts
Argument(s): REALLIST

Description

The counts list of real values gives the frequency or relative probability for each bin in a histogram_bin_uncertain specification. These are paired with the specified abscissas. See histogram_bin_uncertain for details.

7.4.19.5 initial_point

- Keywords Area
- variables
- histogram_bin_uncertain
- initial_point

Initial values for variables

Specification

Alias: none
Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.19.6 descriptors

- **Keywords Area**
- **variables**
- **histogram_bin_uncertain**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** `huv_bin_descriptors`

**Argument(s):** STRINGLIST

**Default:** `hbuv_{i}`

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

### 7.4.20 poisson_uncertain

- **Keywords Area**
- **variables**
- **poisson_uncertain**

Aleatory uncertain discrete variable - Poisson

**Topics**

This keyword is related to the topics:

- **discrete_variables**
- **aleatory_uncertain_variables**
7.4 variables

Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no poisson uncertain variables

**Child Keywords:**
The Poisson distribution is used to predict the number of discrete events that happen in a single time interval. The random events occur uniformly and independently. The expected number of occurrences in a single time interval is $\lambda$, which must be a positive real number. For example, if events occur on average 4 times per year and we are interested in the distribution of events over six months, $\lambda$ would be 2. However, if we were interested in the distribution of events occurring over 5 years, $\lambda$ would be 20.

The probability mass function for the poisson distribution is given by:

$$f(x) = \frac{\lambda^x e^{-\lambda}}{x!}$$

where

- $\lambda$ is the expected number of events occurring in a single time interval
- $x$ is the number of events that occur in this time period
- $f(x)$ is the probability that $x$ events occur in this time period

Theory

When used with some methods such as design of experiments and multidimensional parameter studies, distribution bounds are inferred to be $[0, \mu + 3\sigma]$.

For some methods, including vector and centered parameter studies, an initial point is needed for the uncertain variables. When not given explicitly, these variables are initialized to their means.

7.4.20.1 lambdas

- Keywords Area
  - variables
  - poisson_uncertain
  - lambdas

The parameter for the Poisson distribution, the expected number of events in the time interval of interest

Specification

**Alias**: none

**Argument(s)**: REALLIST
Description

The density function for the Poisson distribution is given by:

\[ f(x) = \frac{\lambda^x e^{-\lambda}}{x!} \]

where \( \lambda \) is the frequency of events happening, and \( x \) is the number of events that occur.

7.4.20.2 initial_point

- Keywords Area
- variables
- poisson_uncertain
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): INTEGERLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.20.3 descriptors

- Keywords Area
- variables
- poisson_uncertain
- descriptors

Labels for the variables
Specification

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** $puv_{i}$

**Description**

The optional variable labels specification *descriptors* is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.21  binomial_uncertain

- Keywords Area
- variables
- binomial_uncertain

Aleatory uncertain discrete variable - binomial

**Topics**

This keyword is related to the topics:

- discrete_variables
- aleatory_uncertain_variables

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** no binomial uncertain variables

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<td>Required</td>
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<tr>
<td>Optional</td>
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</tbody>
</table>

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The binomial distribution describes probabilities associated with a series of independent Bernoulli trials. A Bernoulli trial is an event with two mutually exclusive outcomes, such as 0 or 1, yes or no, success or fail. The probability of success remains the same (the trials are independent).

The density function for the binomial distribution is given by:

\[ f(x) = \binom{n}{x} p^x (1-p)^{n-x} \]

where \( p \) is the probability of failure per trial, \( n \) is the number of trials and \( x \) is the number of successes.

### Theory

The binomial distribution is typically used to predict the number of failures or defective items in a total of \( n \) independent tests or trials, where each trial has the probability \( p \) of failing or being defective.

#### 7.4.21.1 probability_per_trial

- **Keywords Area**
- **variables**
- **binomial_uncertain**
- **probability_per_trial**

A distribution parameter for the binomial distribution

### Specification

**Alias:** prob_per_trial

**Argument(s):** REALLIST

### Description

The binomial distribution is typically used to predict the number of failures (or defective items or some type of event) in a total of \( n \) independent tests or trials, where each trial has the probability \( p \) of failing or being defective. Each particular test can be considered as a Bernoulli trial.

#### 7.4.21.2 num_trials

- **Keywords Area**
- **variables**
- **binomial_uncertain**
- **num_trials**

A distribution parameter
Specification

Alias: none
Argument(s): INTEGERLIST

Description

The binomial distribution is typically used to predict the number of failures (or defective items or some type of event) in a total of \( n \) independent tests or trials, where each trial has the probability \( p \) of failing or being defective. Each particular test can be considered as a Bernoulli trial.

7.4.21.3  initial_point

- Keywords Area
- variables
- binomial_uncertain
- initial_point

Initial values for variables

Specification

Alias: none
Argument(s): INTEGERLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4 variables

7.4.21.4 descriptors

- Keywords Area
- variables
  - binomial_uncertain
- descriptors

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: biuv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.22 negative_binomial_uncertain

- Keywords Area
- variables
  - negative_binomial_uncertain

Aleatory uncertain discrete variable - negative binomial

Topics

This keyword is related to the topics:

- discrete_variables
- aleatory_uncertain_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no negative binomial uncertain variables
Child Keywords:
### Description

The density function for the negative binomial distribution is given by:

\[
f(x) = \binom{n + x - 1}{x} p^n (1 - p)^x\]

where

- \( p \) is the probability of success per trial
- \( n \) is the number of successful trials
- \( X \) is the number of failures

### Theory

The negative binomial distribution is typically used to predict the number of failures observed when repeating a test until a total of \( n \) successes have occurred, where each test has a probability \( p \) of success.

7.4.22.1 probability_per_trial

- Keywords Area
- variables
- negative_binomial_uncertain
- probability_per_trial

A negative binomial distribution parameter

### Specification

**Alias**: prob_per_trial

**Argument(s)**: REALLIST
Description

The negative binomial distribution is typically used to predict the number of failures observed when repeating a test until a total of $n$ successes have occurred, where each test has a probability $p$ of success.

The density function for the negative binomial distribution is given by:

$$f(x) = \binom{n+x-1}{x} p^n (1-p)^x$$

where

- $p$ is the probability of success per trial
- $n$ is the number of successful trials
- $X$ is the number of failures

7.4.22.2 num_trials

- Keywords Area
- variables
- negative_binomial_uncertain
- num_trials

A negative binomial distribution parameter

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

Description

The negative binomial distribution is typically used to predict the number of failures observed when repeating a test until a total of $n$ successes have occurred, where each test has a probability $p$ of success.

The density function for the negative binomial distribution is given by:

$$f(x) = \binom{n+x-1}{x} p^n (1-p)^x$$

where

- $p$ is the probability of success per trial
- $n$ is the number of successful trials
- $X$ is the number of failures

7.4.22.3 initial_point

- Keywords Area
- variables
- negative_binomial_uncertain
- initial_point

Initial values for variables
Specification

Alias: none
Argument(s): INTEGERLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

\[ \text{• Mean/midpoint (or next smaller integer value) when doubly-bounded} \]

\[ \text{• 0 when unbounded} \]

\[ \text{• 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound} \]

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.22.4 descriptors

\[ \text{• Keywords Area} \]
\[ \text{• variables} \]
\[ \text{• negative_binomial_uncertain} \]
\[ \text{• descriptors} \]

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: nbuv_(i)

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.
7.4.23  geometric_uncertain

- Keywords Area
- variables
- geometric_uncertain

Aleatory uncertain discrete variable - geometric

Topics

This keyword is related to the topics:

- discrete_variables
- aleatory_uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no geometric uncertain variables

Child Keywords:

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<tr>
<td>Optional</td>
<td>descriptors</td>
<td></td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

The geometric distribution represents the number of successful trials that might occur before a failure is observed. The density function for the geometric distribution is given by:

\[ f(x) = p(1 - p)^x \]

where \( p \) is the probability of failure per trial.

7.4.23.1  probability_per_trial

- Keywords Area
- variables
- geometric_uncertain
- probability_per_trial

Geometric distribution parameter
### Specification

**Alias:** prob_per_trial  
**Argument(s):** REALLIST

**Description**

The geometric distribution represents the number of successful trials that occur before a failure is observed. The density function for the geometric distribution is given by:

\[ f(x) = p(1 - p)^x \]

where \( p \) is the probability of failure per trial and \( x \) is the number of successful trials.

### 7.4.23.2 initial_point

- **Keywords Area**
- **variables**
- **geometric_uncertain**
- **initial_point**

Initial values for variables

### Specification

**Alias:** none  
**Argument(s):** INTEGERLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4 variables

7.4.23.3 descriptors

- Keywords Area
- variables
- geometric_uncertain
- descriptors

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: geuv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.24 hypergeometric_uncertain

- Keywords Area
- variables
- hypergeometric_uncertain

Aleatory uncertain discrete variable - hypergeometric

Topics

This keyword is related to the topics:

- discrete_variables
- aleatory_uncertain_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no hypergeometric uncertain variables
Child Keywords:
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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>Parameter for the hypergeometric probability distribution describing the size of the total population</td>
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<td>Distribution parameter for the hypergeometric distribution describing the size of the population subset of interest</td>
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<td>Distribution parameter for the hypergeometric distribution describing the number of draws from a combined population</td>
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</tr>
<tr>
<td>Optional</td>
<td></td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

**Description**

The hypergeometric probability density is used when sampling without replacement from a total population of elements where

- The resulting element of each sample can be separated into one of two non-overlapping sets
- The probability of success changes with each sample.

The density function for the hypergeometric distribution is given by:

\[
 f(x) = \binom{m}{x} \binom{N-m}{n-x} / \binom{N}{n}
\]

where the three distribution parameters are:

- \( N \) is the total population
- \( m \) is the number of items in the selected population (e.g. the number of white balls in the full urn of \( N \) items)
- \( n \) is the size of the sample drawn (e.g. number of balls drawn)

In addition,

- \( x \), the abscissa of the density function, indicates the number of successes (e.g. drawing a white ball)
- \( \binom{a}{b} \) indicates a binomial coefficient ("a choose b")
The hypergeometric is often described using an urn model. For example, say we have a total population containing N balls, and we know that m of the balls are white and the remaining balls are green. If we draw n balls from the urn without replacement, the hypergeometric distribution describes the probability of drawing x white balls.

### 7.4.24.1 total_population

- **Keywords Area**
- **variables**
- **hypergeometric_uncertain**
- **total_population**

Parameter for the hypergeometric probability distribution describing the size of the total population

**Specification**

- **Alias**: none
- **Argument(s)**: INTEGERLIST

**Description**

The density function for the hypergeometric distribution is given by:

\[
f(x) = \binom{m}{x} \binom{N-m}{n-x} \binom{N}{n}
\]

where the three distribution parameters are:

- \(N\) is the total population
- \(m\) is the number of items in the selected population (e.g. the number of white balls in the full urn of N items)
- \(n\) is the size of the sample drawn (e.g. number of balls drawn)

In addition,

- \(x\), the abscissa of the density function, indicates the number of successes (e.g. drawing a white ball)
- \(\binom{a}{b}\) indicates a binomial coefficient ("a choose b")

### 7.4.24.2 selected_population

- **Keywords Area**
- **variables**
- **hypergeometric_uncertain**
- **selected_population**

Distribution parameter for the hypergeometric distribution describing the size of the population subset of interest
Specification

**Alias:** none

**Argument(s):** INTEGERLIST

Description

The density function for the hypergeometric distribution is given by:

\[ f(x) = \binom{m}{x} \binom{N-m}{n-x} \binom{N}{n} \]

where the three distribution parameters are:

- N is the total population
- \( m \) is the number of items in the selected population (e.g. the number of white balls in the full urn of N items)
- n is the size of the sample drawn (e.g. number of balls drawn)

In addition,

- x, the abscissa of the density function, indicates the number of successes (e.g. drawing a white ball)
- \( \binom{a}{b} \) indicates a binomial coefficient ("a choose b")

7.4.24.3 num_drawn

- **Keywords Area**
- **variables**
- **hypergeometric_uncertain**
- **num_drawn**

Distribution parameter for the hypergeometric distribution describing the number of draws from a combined population
7.4 variables

- N is the total population
- m is the number of items in the selected population (e.g. the number of white balls in the full urn of N items)
- n is the size of the sample drawn (e.g. number of balls drawn)

In addition,

- x, the abscissa of the density function, indicates the number of successes (e.g. drawing a white ball)
- \( \binom{a}{b} \) indicates a binomial coefficient ("a choose b")

7.4.24.4 initial_point

- Keywords Area
- variables
- hypergeometric_uncertain
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): INTEGERLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4.24.5 descriptors

- Keywords Area
- variables
- hypergeometric_uncertain
- descriptors

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: hguv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.25 histogram_point_uncertain

- Keywords Area
- variables
- histogram_point_uncertain

Aleatory uncertain variable - discrete histogram

Topics

This keyword is related to the topics:

- discrete_variables
- aleatory_uncertain_variables

Specification

Alias: none
Argument(s): none
Default: no histogram point uncertain variables

Child Keywords:
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<td>String (categorical) valued point histogram variable</td>
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**Description**

Histogram uncertain variables are typically used to model a set of empirical data. When the variables take on only discrete values or categories, a discrete, or point histogram is used to describe their probability mass function (one could think of this as a histogram_bin_uncertain variable with "bins" of zero width). Dakota supports integer-, string-, and real-valued point histograms.

Point histograms are similar to discrete_design_set and discrete_state_set, but as they are uncertain variables, include the relative probabilities of observing the different values within the set.

The histogram_point_uncertain keyword is followed by one or more of integer, string, or real, each of which specify the number of variables to be characterized as discrete histograms of that sub-type.

Each discrete histogram variable is specified by one or more abscissa/ count pairs. The abscissas, are the possible values the variable can take on ("x" coordinates of type integer, string, or real), and must be specified in increasing order. These are paired with counts which provide the frequency of the given value or string, relative to other possible values/strings.

Thus, to fully specify a point-based histogram with n points, n (x,c) pairs must be specified with the following features:

- x is the point value (integer, string, or real) and c is the corresponding count for that value.
- the x values must be strictly increasing (lexicographically for strings).
- all c values must be positive.
- a minimum of one pair must be specified for each point-based histogram.

**Examples**

The pairs_per_variable specification provides for the proper association of multiple sets of (x,c) or (x,y) pairs with individual histogram variables. For example, in the following specification,

```plaintext
histogram_point_uncertain
    integer = 2
    pairs_per_variable = 2 3
    abscissas = 3 4 100 200 300
    counts = 1 1 1 2 1
```

pairs_per_variable associates the (x,c) pairs ((3,1),(4,1)) with one point-based histogram variable (where the values 3 and 4 are equally probable) and associates the (x,c) pairs ((100,1),(200,2),(300,1)) with a second point-based histogram variable (where the value 200 is twice as probable as either 100 or 300).
See Also

These keywords may also be of interest:

- histogram_bin_uncertain

FAQ

**Difference between bin and point histograms:** A (continuous) bin histogram specifies bins of non-zero width, whereas a (discrete) point histogram specifies individual point values, which can be thought of as bins with zero width. In the terminology of LHS\cite{93}, the bin pairs specification defines a "continuous linear" distribution and the point pairs specification defines a "discrete histogram" distribution (although the points are real-valued, the number of possible values is finite).

7.4.25.1 integer

- Keywords Area
- variables
- histogram_point_uncertain
- integer

Integer valued point histogram variable

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

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<td>Counts for integer-valued point histogram</td>
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**7.4 variables**

<table>
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<th>descriptors</th>
<th>Labels for the variables</th>
</tr>
</thead>
</table>

**Description**

This probability mass function is integer-valued; the abscissa values must all be integers. The \( n \) abscissa values are paired with \( n \) counts which indicate the relative frequency (mass) of each integer relative to the other specified integers.

**Examples**

```plaintext
histogram_point_uncertain
  integer = 2
  pairs_per_variable = 2 3
  abscissas = 3 4 100 200 300
  counts = 1 1 1 2 1
```

There are two variables, the first one has two possible integer values which are equally probable. The second one has three options, and 200 is twice as probable as either 100 or 300.

7.4.25.1.1 pairs_per_variable

- **Keywords Area**
- variables
- histogram_point_uncertain
- integer
- pairs_per_variable

Number of pairs defining each histogram point integer variable

**Specification**

*Alias:* num_pairs

*Argument(s):* INTEGERLIST

*Default:* equal distribution

**Description**

By default, the list of abscissas and counts will be evenly divided among the histogram point integer variables. The number of pairs_per_variable specifies the apportionment of abscissa/count pairs among the histogram point integer variables. It must specify one integer \( \geq 1 \) per variable that indicates how many of the \((\text{abscissa}, \text{count}) = (x,c)\) pairs to associate with that variable.

7.4.25.1.2 abscissas

- **Keywords Area**
- variables
- histogram_point_uncertain
- integer
• **abscissas**

Integer abscissas for a point histogram

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

A list of integer abscissa ("x" coordinate) values characterizing the probability density function for each of the integer histogram_point_uncertain variables. These must be listed in increasing order for each variable, and are paired with counts. See histogram_point_uncertain for details and examples.

7.4.25.1.3 **counts**

• **Keywords Area**

• variables

• histogram_point_uncertain

• integer

• counts

Counts for integer-valued point histogram

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Description**

Count or frequency for each of abscissas. See histogram_point_uncertain for details and examples.

7.4.25.1.4 **initial_point**

• **Keywords Area**

• variables

• histogram_point_uncertain

• integer

• initial_point

Initial values for variables

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST
7.4 variables

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.25.1.5 descriptors

- Keywords Area
- variables
- histogram_point_uncertain
- integer
- descriptors

Labels for the variables

Specification

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** hpi\(_v\)_\(_i\)

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.25.2 string

- Keywords Area
- variables
- histogram_point_uncertain
- string

String (categorical) valued point histogram variable
Specification

Alias: none

Arguments: INTEGER

Child Keywords:

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</tr>
</tbody>
</table>

Description

This probability mass function is string-valued; the abscissa values must all be strings. The \( n \) abscissa values are paired with \( n \) counts which indicate the relative frequency (mass) of each string relative to the other specified strings.

Examples

```
histogram_point_uncertain
string = 2
pairs_per_variable = 2 3
abscissas = 'no' 'yes' 'function1' 'function2' 'function3'
counts = 1 1 1 2 1
descriptors = 'vote' 'which_function'
```

Here there are two variables, the first one ('vote') has two possible string values 'yes' and 'no' which are equally probable. The second one has three options for 'which_function', and 'function2' is twice as probable as 'function1' or 'function3'.

7.4.25.2.1 pairs_per_variable

- Keywords Area
- variables
- histogram_point_uncertain
- string
- pairs_per_variable

Number of pairs defining each histogram point string variable
7.4 variables

Specification

**Alias:** num_pairs

**Argument(s):** INTEGERLIST

**Default:** equal distribution

**Description**

By default, the list of abscissas and counts will be evenly divided among the histogram point string variables. The number of pairs_per_variable specifies the apportionment of abscissa/count pairs among the histogram point string variables. It must specify one integer $\geq 1$ per variable that indicates how many of the (abscissa, count) = (x,c) pairs to associate with that variable.

7.4.25.2.2 abscissas

- **Keywords Area**
- variables
- histogram_point_uncertain
- string
- abscissas

String abscissas for a point histogram

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Description**

A list of string abscissa ("x" coordinate) values characterizing the probability density function for each of the string histogram_point_uncertain variables. These must be listed in (lexicographically) increasing order for each variable, and are paired with counts. See histogram_point_uncertain for details and examples.

7.4.25.2.3 counts

- **Keywords Area**
- variables
- histogram_point_uncertain
- string
- counts

Counts for string-valued point histogram

**Specification**

**Alias:** none

**Argument(s):** REALLIST
Description

Count or frequency for each of abscissas. See histogram_point_uncertain for details and examples.

7.4.25.2.4 initial_point

- Keywords Area
- variables
- histogram_point_uncertain
- string
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): STRINGLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.25.2.5 descriptors

- Keywords Area
- variables
- histogram_point_uncertain
- string
- descriptors

Labels for the variables
7.4 variables

Specification

 Alias: none
 Argument(s): STRINGLIST
 Default: hpsv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.25.3 real

• Keywords Area
  • variables
  • histogram_point_uncertain
  • real

Real valued point histogram variable

Specification

 Alias: none
 Argument(s): INTEGER

Child Keywords:

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<tr>
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<td>descriptors</td>
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<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

This probability mass function is real-valued; the abscissa values must all be integers. The $n$ abscissa values are paired with $n$ counts which indicate the relative frequency (mass) of each real relative to the other specified reals.
Examples

```plaintext
histogram_point_uncertain
  real = 2
  pairs_per_variable = 2 3
  abscissas = 3.1415 4.5389 100 200.112345 300
  counts = 1 1 1 2 1
```

There are two variables, the first one has two possible real values which are equally probable. The second one has three possible real value options, and 200.112345 is twice as probable as either 100 or 300.

7.4.25.3.1 pairs_per_variable

- **Keywords Area**
- **variables**
- **histogram_point_uncertain**
- **real**
- **pairs_per_variable**

Number of pairs defining each histogram point real variable

**Specification**

**Alias:** num_pairs

**Argument(s):** INTEGERLIST

**Default:** equal distribution

**Description**

By default, the list of abscissas and counts will be evenly divided among the histogram point real variables. The number of pairs_per_variable specifies the apportionment of abscissa/count pairs among the histogram point real variables. It must specify one integer \( \geq 1 \) per variable that indicates how many of the \((\text{abscissa, count}) = (x, c)\) pairs to associate with that variable.

7.4.25.3.2 abscissas

- **Keywords Area**
- **variables**
- **histogram_point_uncertain**
- **real**
- **abscissas**

Real abscissas for a point histogram

**Specification**

**Alias:** none

**Argument(s):** REALLIST
Description

A list of real abscissa ("x" coordinate) values characterizing the probability density function for each of the real histogram_point_uncertain variables. These must be listed in increasing order for each variable, and are paired with counts. See histogram_point_uncertain for details and examples.

7.4.25.3.3 counts

- Keywords Area
- variables
- histogram_point_uncertain
- real
- counts

Counts for real-valued point histogram

Specification

Alias: none

Argument(s): REALLIST

Description

Count or frequency for each of abscissas. See histogram_point_uncertain for details and examples.

7.4.25.3.4 initial_point

- Keywords Area
- variables
- histogram_point_uncertain
- real
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:
• Mean/midpoint (or next smaller integer value) when doubly-bounded

• 0 when unbounded

• 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.25.3.5 descriptors

• Keywords Area
• variables
• histogram_point_uncertain
• real
• descriptors

Labels for the variables

Specification

Alias: none

Argument(s): STRINGLIST

Default: hpruv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.26 uncertain_correlation_matrix

• Keywords Area
• variables
• uncertain_correlation_matrix

Correlation among aleatory uncertain variables

Specification

Alias: none

Argument(s): REALLIST

Default: identity matrix (uncorrelated)
Description

Aleatory uncertain variables may have correlations specified through use of an `uncertain_correlation_matrix` specification. This specification is generalized in the sense that its specific meaning depends on the nondeterministic method in use.

When the method is a nondeterministic sampling method (i.e., `sampling`), then the correlation matrix specifies rank correlations [53].

When the method is a reliability (i.e., `local_reliability` or `global_reliability`) or stochastic expansion (i.e., `polynomial_chaos` or `stoch_collocation`) method, then the correlation matrix specifies correlation coefficients (normalized covariance) [43].

In either of these cases, specifying the identity matrix results in uncorrelated uncertain variables (the default). The matrix input should be symmetric and have all $n^2$ entries where $n$ is the total number of aleatory uncertain variables.

Ordering of the aleatory uncertain variables is:

1. normal
2. lognormal
3. uniform
4. loguniform
5. triangular
6. exponential
7. beta
8. gamma
9. gumbel
10. frechet
11. weibull
12. histogram bin
13. poisson
14. binomial
15. negative binomial
16. geometric
17. hypergeometric
18. histogram point

When additional variable types are activated, they assume uniform distributions, and the ordering is as listed on variables.

Examples

Consider the following random variables, distributions and correlations:

- $X_1$, normal, uncorrelated with others
- $X_2$, normal, correlated with $X_3$, $X_4$ and $X_5$
- $X_3$, weibull, correlated with $X_5$
• $X_3$, exponential, correlated with $X_3$, $X_4$ and $X_5$

• $X_5$, normal, correlated with $X_3$ These correlations are captured by the following commands (order of the variables is respected).

`uncertain_correlation_matrix`

```text
# ordering normal, exponential, weibull
1.00 0.00 0.00 0.00 0.00
0.00 1.00 0.50 0.24 0.78
0.00 0.50 1.00 0.00 0.20
0.00 0.24 0.00 1.00 0.49
0.00 0.78 0.20 0.49 1.0
```

7.4.27 continuous_interval_uncertain

• Keywords Area

• variables

• continuous_interval_uncertain

Epistemic uncertain variable - values from one or more continuous intervals

Topics

This keyword is related to the topics:

• continuous_variables

• epistemic_uncertain_variables

Specification

Alias: interval_uncertain

Argument(s): INTEGER

Default: no continuous interval uncertain variables

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>num_intervals</td>
<td></td>
<td>Specify the number of intervals for each variable</td>
</tr>
<tr>
<td>Optional</td>
<td>interval_probabilities</td>
<td></td>
<td>Assign probability mass to each interval</td>
</tr>
<tr>
<td>Required</td>
<td>lower_bounds</td>
<td></td>
<td>Specify minimum values</td>
</tr>
<tr>
<td>Required</td>
<td>upper_bounds</td>
<td></td>
<td>Specify maximum values</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_point</td>
<td></td>
<td>Initial values for variables</td>
</tr>
</tbody>
</table>
**Description**

Continuous interval uncertain variables are epistemic types. They can specify a single interval per variable which may be used in interval analysis, where the goal is to determine the interval bounds on the output corresponding to the interval bounds on the input. All values between the bounds are permissible. More detailed continuous interval representations can specify a set of belief structures based on intervals that may be contiguous, overlapping, or disjoint. This is used in specifying the inputs necessary for an epistemic uncertainty analysis using Dempster-Shafer theory of evidence.

Other epistemic types include:

- `discrete_interval_uncertain`
- `discrete_uncertain_set integer`
- `discrete_uncertain_set string`
- `discrete_uncertain_set real`

**Examples**

The following specification is for an interval analysis:

```plaintext
continuous_interval_uncertain = 2
lower_bounds = 2.0 4.0
upper_bounds = 2.5 5.0
```

The following specification is for a Dempster-Shafer analysis:

```plaintext
continuous_interval_uncertain = 2
num_intervals = 3 2
interval_probs = 0.25 0.5 0.25 0.4 0.6
lower_bounds = 2.0 4.0 4.5 1.0 3.0
upper_bounds = 2.5 5.0 6.0 5.0 5.0
```

Here there are 2 interval uncertain variables. The first one is defined by three intervals, and the second by two intervals. The three intervals for the first variable have basic probability assignments of 0.2, 0.5, and 0.3, respectively, while the basic probability assignments for the two intervals for the second variable are 0.4 and 0.6. The basic probability assignments for each interval variable must sum to one. The interval bounds for the first variable are [2, 2.5], [4, 5], and [4.5, 6], and the interval bounds for the second variable are [1.0, 5.0] and [3.0, 5.0]. Note that the intervals can be overlapping or disjoint. The BPA for the first variable indicates that it is twice as likely that the value occurs on the interval [4,5] than either [2,2,5] or [4,5,6].

**Theory**

The continuous interval uncertain variable is NOT a probability distribution. Although it may seem similar to a histogram, the interpretation of this uncertain variable is different. It is used in epistemic uncertainty analysis, where one is trying to model uncertainty due to lack of knowledge. The continuous interval uncertain variable is used in both interval analysis and in Dempster-Shafer theory of evidence.

- interval analysis -only one interval is allowed for each `continuous_interval_uncertain` variable
- the interval is defined by lower and upper bounds
- the value of the random variable lies somewhere in this interval
- output is the minimum and maximum function value conditional on the specified interval
Dempster-Shafer theory of evidence - multiple intervals can be assigned to each continuous_interval_uncertain variable - a Basic Probability Assignment (BPA) is associated with each interval. The BPA represents a probability that the value of the uncertain variable is located within that interval. Each interval is defined by lower and upper bounds - outputs are called "belief" and "plausibility." Belief represents the smallest possible probability that is consistent with the evidence, while plausibility represents the largest possible probability that is consistent with the evidence. Evidence is the intervals together with their BPA.

7.4.27.1 num_intervals

- Keywords Area

- variables

- continuous_interval_uncertain

- num_intervals

Specify the number of intervals for each variable

Specification

Alias: iuv_num_intervals

Argument(s): INTEGERLIST

Default: Equal apportionment of intervals among variables

Description

In Dakota, epistemic uncertainty analysis is performed using either interval estimation or Dempster-Shafer theory of evidence. In these approaches, one does not assign a probability distribution to each uncertain input variable. Rather, one divides each uncertain input variable into one or more intervals. The input parameters are only known to occur within intervals; nothing more is assumed. num_intervals specifies the number of such intervals associated with each interval uncertain parameter.

7.4.27.2 interval_probabilities

- Keywords Area

- variables

- continuous_interval_uncertain

- interval_probabilities

Assign probability mass to each interval

Specification

Alias: interval_probs iuv_interval_probs

Argument(s): REALLIST

Default: Equal probability assignments for each interval (1/num_intervals[i])
Description

The basic probability assignments for each interval variable must sum to one. For example, if an interval variable is defined with three intervals, the probabilities for these intervals could be 0.2, 0.5, and 0.3 which sum to one, but could not be 0.5, 0.5, and 0.5 which do not sum to one.

7.4.27.3 lower_bounds

- Keywords Area
- variables
- continuous_interval_uncertain
- lower_bounds

Specify minimum values

Specification

Alias: none
Argument(s): REALLIST

Description

Specify minimum values

7.4.27.4 upper_bounds

- Keywords Area
- variables
- continuous_interval_uncertain
- upper_bounds

Specify maximum values

Specification

Alias: none
Argument(s): REALLIST

Description

Specify maximum values
7.4.27.5 initial_point

- Keywords Area
- variables
  - continuous_interval_uncertain
  - initial_point

Initial values for variables

**Specification**

**Alias**: none

**Argument(s)**: REALLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.27.6 descriptors

- Keywords Area
- variables
  - continuous_interval_uncertain
  - descriptors

Labels for the variables

**Specification**

**Alias**: iuv_descriptors

**Argument(s)**: STRINGLIST

**Default**: ciuv_[i]
Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.28 discrete_interval_uncertain

- Keywords Area
- variables
- discrete_interval_uncertain

Epistemic uncertain variable - values from one or more discrete intervals

Topics

This keyword is related to the topics:

- discrete_variables
- epistemic_uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: No discrete interval uncertain variables

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakoita Keyword Description</th>
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<td>Assign probability mass to each interval</td>
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<td>Specify maximum values</td>
</tr>
<tr>
<td>Optional</td>
<td>initial_point</td>
<td>initial_point</td>
<td>Initial values for variables</td>
</tr>
<tr>
<td>Optional</td>
<td>descriptors</td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

Description

Discrete interval uncertain variables are epistemic types. They can specify a single interval per variable which may be used in interval analysis, where the goal is to determine the interval bounds on the output corresponding to the
interval bounds on the input. Permissible values are any integer within the bound. More detailed continuous interval representations can specify a set of belief structures based on intervals that may be contiguous, overlapping, or disjoint. This is used in specifying the inputs necessary for an epistemic uncertainty analysis using Dempster-Shafer theory of evidence.

Other epistemic types include:

- `continuous_interval_uncertain`
- `discrete_uncertain_set integer`
- `discrete_uncertain_set string`
- `discrete_uncertain_set real`

Examples

Let $d1$ be 2, 3 or 4 with probability 0.2, 4 or 5 with probability 0.5 and 6 with probability 0.3. Let $d2$ be 4, 5 or 6 with probability 0.4 and 6, 7 or 8 with probability 0.6. The following specification is for a Dempster-Shafer analysis:

```plaintext
discrete_interval_uncertain = 2
num_intervals = 3 2
interval_probs = 0.2 0.5 0.3 0.4 0.6
lower_bounds = 2 4 6 4 6
upper_bounds = 4 5 6 6 8
```

Theory

- Dempster-Shafer theory of evidence - multiple intervals can be assigned to each `discrete_interval_uncertain` variable - a Basic Probability Assignment (BPA) is associated with each interval. The BPA represents a probability that the value of the uncertain variable is located within that interval. - each interval is defined by lower and upper bounds - outputs are called “belief” and “plausibility.” Belief represents the smallest possible probability that is consistent with the evidence, while plausibility represents the largest possible probability that is consistent with the evidence. Evidence is the intervals together with their BPA.

7.4.28.1 `num_intervals`

- `Keywords Area`
- `variables`
- `discrete_interval_uncertain`
- `num_intervals`

Specify the number of intervals for each variable

Specification

- **Alias:** none
- **Argument(s):** INTEGERLIST
- **Default:** Equal apportionment of intervals among variables
Description

In Dakota, epistemic uncertainty analysis is performed using either interval estimation or Dempster-Shafer theory of evidence. In these approaches, one does not assign a probability distribution to each uncertain input variable. Rather, one divides each uncertain input variable into one or more intervals. The input parameters are only known to occur within intervals; nothing more is assumed. num_intervals specifies the number of such intervals associated with each interval uncertain parameter.

7.4.28.2 interval_probabilities

- Keywords Area
- variables
- discrete_interval_uncertain
- interval_probabilities

Assign probability mass to each interval

Specification

Alias: interval_probs range_probabilities range_probs

Argument(s): REALLIST

Default: Equal probability assignments for each interval (1/num_intervals[i])

Description

The basic probability assignments for each interval variable must sum to one. For example, if an interval variable is defined with three intervals, the probabilities for these intervals could be 0.2, 0.5, and 0.3 which sum to one, but could not be 0.5,0.5, and 0.5 which do not sum to one.

7.4.28.3 lower_bounds

- Keywords Area
- variables
- discrete_interval_uncertain
- lower_bounds

Specify minimum values

Specification

Alias: none

Argument(s): INTEGERLIST

Description

Specify minimum values
7.4.28.4 upper_bounds

- Keywords Area
- variables
- discrete_interval_uncertain
- upper_bounds

Specify maximum values

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Specify maximum values

7.4.28.5 initial_point

- Keywords Area
- variables
- discrete_interval_uncertain
- initial_point

Initial values for variables

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.28 descriptors

- **Keywords Area**
- **variables**
- **discrete_interval_uncertain**
- **descriptors**

Labels for the variables

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** `diuv_{i}`

**Description**

The optional variable labels specification `descriptors` is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

### 7.4.29 discrete_uncertain_set

- **Keywords Area**
- **variables**
- **discrete_uncertain_set**

Epistemic uncertain variable - discrete set-valued

**Topics**

This keyword is related to the topics:

- **discrete_variables**
- **epistemic_uncertain_variables**

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**
### Description

Discrete uncertain variables whose values come from a set of admissible elements. Each variable specified must be of type integer, string, or real.

#### 7.4.29.1 integer

- **Keywords Area**
  - variables
  - discrete_uncertain_set
  - integer

Discrete, epistemic uncertain variable - integers within a set

### Topics

This keyword is related to the topics:

- discrete_variables
- epistemic_uncertain_variables

### Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no discrete uncertain set integer variables

**Child Keywords:**
Description

Discrete set variables may be used to specify categorical choices which are epistemic. For example, if we have three possible forms for a physics model (model 1, 2, or 3) and there is epistemic uncertainty about which one is correct, a discrete uncertain set may be used to represent this type of uncertainty.

This variable is defined by a set of integers, in which the discrete value may take any value within the integer set (for example, the set may be defined as 1, 2, and 4).

Other epistemic types include:

- `continuous_interval_uncertain`
- `discrete_interval_uncertain`
- `discrete_uncertain_set string`
- `discrete_uncertain_set real`

Examples

Let `d1` be 2 or 13 and `d2` be 4, 5 or 26. The following specification is for an interval analysis:

```
discrete_uncertain_set
integer
num_set_values 2 3
set_values 2 13 4 5 26
descriptors 'di1' 'di2'
```
The discrete uncertain set-integer variable is NOT a discrete random variable. It can be contrasted to the histogram-defined random variables: histogram_bin_uncertain and histogram_point_uncertain. It is used in epistemic uncertainty analysis, where one is trying to model uncertainty due to lack of knowledge.

The discrete uncertain set integer variable is used in both interval analysis and in Dempster-Shafer theory of evidence.

- **interval analysis** -the values are integers, equally weighted -the true value of the random variable is one of the integers in this set -output is the minimum and maximum function value conditional on the specified inputs

- **Dempster-Shafer theory of evidence** -the values are integers, but they can be assigned different weights -outputs are called "belief" and "plausibility." Belief represents the smallest possible probability that is consistent with the evidence, while plausibility represents the largest possible probability that is consistent with the evidence. Evidence is the values together with their weights.

### 7.4.29.1.1 elements_per_variable

- **Keywords Area**
- **variables**
- **discrete_uncertain_set**
- **integer**
- **elements_per_variable**

Number of admissible elements for each set variable

### Specification

**Alias:** num_set_values

**Argument(s):** INTEGERLIST

**Default:** equal distribution

### Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

### 7.4.29.1.2 elements

- **Keywords Area**
- **variables**
- **discrete_uncertain_set**
- **integer**
- **elements**

The permissible values for each discrete variable
7.4 variables

Specification

Alias: set_values
Argument(s): INTEGERLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.29.1.3 set_probabilities

• Keywords Area
• variables
• discrete_uncertain_set
• integer
• set_probabilities

This keyword defines the probabilities for the various elements of discrete sets.

Specification

Alias: set_probs
Argument(s): REALLIST
Default: Equal probability assignments for each set member (1/num_set_values[i])

Description

There are three types of discrete_uncertain_set variables: integer, string, or real sets. With each of these types, one defines the number of elements of the set per that variable, the values of those elements, and the associated probabilities. For example, if one has an integer discrete uncertain set variable with 3 elements {3,4,8}, then one could define the probabilities associated with those set elements as (for example) 0.2, 0.5, and 0.3. The set_probabilities for a particular variable should sum to one over all the elements in that set.

7.4.29.1.4 categorical

• Keywords Area
• variables
• discrete_uncertain_set
• integer
• categorical

Whether the set-valued variables are categorical or relaxable

Specification

Alias: none
Argument(s): STRINGLIST
**Description**

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include ‘yes’, ‘no’, ‘true’, and ‘false’, or any abbreviation in [yYnNtTfF][.]∗

**Examples**

Discrete_design_set variable, ‘rotor_blades’, can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and ‘rotor_blades’ can take on any value between 2 and 7, e.g., 3, 6, or 5.5.

```plaintext
discrete_design_set
  integer 1
  elements 2 4 7
  descriptor ‘rotor_blades’
  categorical ‘no’
```

**7.4.29.1.5 initial_point**

- **Keywords Area**
  - variables
    - discrete_uncertain_set
  - integer
  - initial_point

Initial values for variables

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4.29.1.6 descriptors

- Keywords Area
- variables
- discrete_uncertain_set
- integer
- descriptors

Labels for the variables

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** dusiv_{i}

**Description**

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.29.2 string

- Keywords Area
- variables
- discrete_uncertain_set
- string

Discrete, epistemic uncertain variable - strings within a set

**Topics**

This keyword is related to the topics:

- discrete_variables
- epistemic_uncertain_variables

**Specification**

**Alias:** none

**Argument(s):** INTEGER

**Default:** no discrete uncertain set string variables

**Child Keywords:**
### Description

Discrete set variables may be used to specify categorical choices which are epistemic. For example, if we have three possible forms for a physics model (model 1, 2, or 3) and there is epistemic uncertainty about which one is correct, a discrete uncertain set may be used to represent this type of uncertainty.

This variable is defined by a set of strings, in which the discrete value may take any value within the string set (for example, the set may be defined as 'coarse', 'medium', and 'fine')

Other epistemic types include:

- `continuous_interval_uncertain`
- `discrete_interval_uncertain`
- `discrete_uncertain_set integer`
- `discrete_uncertain_set real`

### Examples

```dakota
discrete_uncertain_set
string
num_set_values 2 3
set_values 'red' 'blue' 'coarse' 'medium' 'fine'
descriptors 'ds1' 'ds2'
```

7.4.29.2.1 `elements_per_variable`

- Keywords Area
- variables
- discrete_uncertain_set
- string
- elements_per_variable

Number of admissible elements for each set variable
7.4 variables

Specification

Alias: num_set_values
Argument(s): INTEGERLIST
Default: equal distribution

Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

7.4.29.2.2 elements

- Keywords Area
- variables
- discrete_uncertain_set
- string
- elements

The permissible values for each discrete variable

Specification

Alias: set_values
Argument(s): STRINGLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.29.2.3 set_probabilities

- Keywords Area
- variables
- discrete_uncertain_set
- string
- set_probabilities

This keyword defines the probabilities for the various elements of discrete sets.

Specification

Alias: set_probs
Argument(s): REALLIST
Default: Equal probability assignments for each set member (1/num_set_values[i])
Description

There are three types of `discrete_uncertain_set` variables: integer, string, or real sets. With each of these types, one defines the number of elements of the set per that variable, the values of those elements, and the associated probabilities. For example, if one has an integer discrete uncertain set variable with 3 elements \{3,4,8\}, then one could define the probabilities associated with those set elements as (for example) 0.2, 0.5, and 0.3. The set_probabilities for a particular variable should sum to one over all the elements in that set.

7.4.29.2.4 initial_point

- Keywords Area
- variables
- discrete_uncertain_set
- string
- initial_point

Initial values for variables

Specification

Alias: none

Argument(s): STRINGLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.29.2.5 descriptors

- Keywords Area
- variables
- discrete_uncertain_set
- string
- descriptors

Labels for the variables
7.4 variables

Specification

Alias: none

Argument(s): STRINGLIST

Default: dussv_{i[1]}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.29.3 real

- Keywords Area
  - variables
    - discrete_uncertain_set
    - real

Discrete, epistemic uncertain variable - real numbers within a set

Topics

This keyword is related to the topics:

- discrete_variables
- epistemic_uncertain_variables

Specification

Alias: none

Argument(s): INTEGER

Default: no discrete uncertain set real variables

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>elements_per_variable</td>
<td>Number of admissible elements for each set variable</td>
</tr>
</tbody>
</table>
### Description

Discrete set variables may be used to specify categorical choices which are epistemic. For example, if we have three possible forms for a physics model (model 1, 2, or 3) and there is epistemic uncertainty about which one is correct, a discrete uncertain set may be used to represent this type of uncertainty.

This variable is defined by a set of reals, in which the discrete variable may take any value defined within the real set (for example, a parameter may have two allowable real values, 3.285 or 4.79).

Other epistemic types include:

- `continuous_interval_uncertain`
- `discrete_interval_uncertain`
- `discrete_uncertain_set integer`
- `discrete_uncertain_set string`

### Examples

Let $d_1$ be 2.1 or 1.3 and $d_2$ be 0.4, 5 or 2.6. The following specification is for an interval analysis:

```plaintext
discrete_uncertain_set
    integer
    num_set_values 2 3
    set_values 2.1 1.3 0.4 5 2.6
    descriptors 'd1' 'd2'
```

### Theory

The `discrete_uncertain_set integer` variable is **NOT** a discrete random variable. It can be contrasted to a the histogram-defined random variables: `histogram_bin_uncertain` and `histogram_point_uncertain`. It is used in epistemic uncertainty analysis, where one is trying to model uncertainty due to lack of knowledge.

The discrete uncertain set integer variable is used in both interval analysis and in Dempster-Shafer theory of evidence.
• interval analysis -the values are integers, equally weighted -the true value of the random variable is one of the integers in this set -output is the minimum and maximum function value conditional on the specified inputs

• Dempster-Shafer theory of evidence -the values are integers, but they can be assigned different weights -outputs are called "belief" and "plausibility." Belief represents the smallest possible probability that is consistent with the evidence, while plausibility represents the largest possible probability that is consistent with the evidence. Evidence is the values together with their weights.

7.4.29.3.1 elements_per_variable

• Keywords Area
• variables
• discrete_uncertain_set
• real
• elements_per_variable

Number of admissible elements for each set variable

Specification

Alias: num_set_values
Argument(s): INTEGERLIST
Default: equal distribution

Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

7.4.29.3.2 elements

• Keywords Area
• variables
• discrete_uncertain_set
• real
• elements

The permissible values for each discrete variable

Specification

Alias: set_values
Argument(s): REALLIST
Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.29.3.3 set_probabilities

- Keywords Area
- variables
- discrete_uncertain_set
- real
- set_probabilities

This keyword defines the probabilities for the various elements of discrete sets.

Specification

**Alias:** set_probs

**Argument(s):** REALIST

**Default:** Equal probability assignments for each set member (1/num_set_values[i])

Description

There are three types of discrete_uncertain_set variables: integer, string, or real sets. With each of these types, one defines the number of elements of the set per that variable, the values of those elements, and the associated probabilities. For example, if one has an integer discrete uncertain set variable with 3 elements {3,4,8}, then one could define the probabilities associated with those set elements as (for example) 0.2, 0.5, and 0.3. The set_probabilities for a particular variable should sum to one over all the elements in that set.

7.4.29.3.4 categorical

- Keywords Area
- variables
- discrete_uncertain_set
- real
- categorical

Whether the set-valued variables are categorical or relaxable

Specification

**Alias:** none

**Argument(s):** STRINGLIST

Description

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include ‘yes’, ‘no’, ‘true’, and ‘false’, or any abbreviation in [yYnNtTfF]\*.
Examples

Discrete_design_set variable, ‘rotor_blades’, can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and ‘rotor_blades’ can take on any value between 2 and 7, e.g., 3, 6, or 5.5.

discrete_design_set
    integer 1
    elements 2 4 7
    descriptor 'rotor_blades'
    categorical 'no'

7.4.29.3.5 initial_point

- Keywords Area
- variables
  - discrete_uncertain_set
  - real
  - initial_point

Initial values for variables

Specification

Alias: none

Argument(s): REALLIST

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
7.4.29.3.6 descriptors

- Keywords Area
- variables
- discrete_uncertain_set
- real
- descriptors

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: dusrv_(i)

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.
The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.30 continuous_state

- Keywords Area
- variables
- continuous_state

State variable - continuous

Topics

This keyword is related to the topics:

- state_variables
- continuous_variables

Specification

Alias: none
Argument(s): INTEGER
Default: No continuous state variables
Child Keywords:
### Description

Continuous state variables are defined by bounds.

Default behavior for most methods is that only the initial_state values are used.

See the [state_variables](#) page for details on the behavior of state variables.

#### 7.4.30.1 initial_state

- **Keywords Area**
- **variables**
- **continuous_state**
- **initial_state**

Initial values for variables

#### Specification

**Alias:** csv_initial_state

**Argument(s):** REALLIST

**Default:** 0.0

#### Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including `continuous_design`, `continuous_state`, `discrete_design_range`, and `discrete_state_range` are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

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Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.30.2 lower_bounds

- **Keywords Area**
- **variables**
- **continuous_state**
- **lower_bounds**

Specify minimum values

**Specification**

**Alias:** `csv_lower_bounds`  
**Argument(s):** `REALLIST`  
**Default:** `-infinity`

**Description**

Specify minimum values

### 7.4.30.3 upper_bounds

- **Keywords Area**
- **variables**
- **continuous_state**
- **upper_bounds**

Specify maximum values

**Specification**

**Alias:** `csv_upper_bounds`  
**Argument(s):** `REALLIST`  
**Default:** `infinity`

**Description**

Specify maximum values
7.4 variables

7.4.30.4 descriptors

- Keywords Area
- variables
- continuous_state
- descriptors

Labels for the variables

Specification

Alias: csv_descriptors
Argument(s): STRINGLIST
Default: csv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.31 discrete_state_range

- Keywords Area
- variables
- discrete_state_range

State variables - discrete range-valued

Topics

This keyword is related to the topics:

- discrete_variables
- state_variables

Specification

Alias: none
Argument(s): INTEGER
Default: No discrete state variables
Child Keywords:
### Required/Optional Group

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<th>Dakota Keyword</th>
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<td>Labels for the variables</td>
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### Description

Discrete state variables defined by bounds (an integer interval).

The details of how to specify this discrete variable are located on the [discrete_variables](#) page.

See the [state_variables](#) page for details on the behavior of state variables.

#### 7.4.31.1 initial_state

- **Keywords Area**
- **variables**
- **discrete_state_range**
- **initial_state**

Initial values for variables

### Specification

**Alias:** dsv_initial_state

**Argument(s):** INTEGERLIST

**Default:** 0

### Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

### Default Behavior

Range variables, including **continuous_design**, **continuous_state**, **discrete_design_range**, and **discrete_state_range** are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound
Set variables, including `discrete_design_set` and `discrete_state_set` default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

### 7.4.3.1.2 lower_bounds

- **Keywords Area**
- **variables**
- **discrete_state_range**
- **lower_bounds**

Specify minimum values

#### Specification

**Alias:** `dsv_lower_bounds`

**Argument(s):** `INTEGERLIST`

**Default:** `INT_MIN`

**Description**

Specify minimum values

### 7.4.3.1.3 upper_bounds

- **Keywords Area**
- **variables**
- **discrete_state_range**
- **upper_bounds**

Specify maximum values

#### Specification

**Alias:** `dsv_upper_bounds`

**Argument(s):** `INTEGERLIST`

**Default:** `INT_MAX`

**Description**

Specify maximum values
7.4.31.4 descriptors

- Keywords Area
- variables
- discrete_state_range
- descriptors

Labels for the variables

Specification

**Alias:** dsv_descriptors

**Argument(s):** STRINGLIST

**Default:** dsriv_{i}

**Description**

The optional variable labels specification *descriptors* is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.32 discrete_state_set

- Keywords Area
- variables
- discrete_state_set

State variable - discrete set-valued

**Topics**

This keyword is related to the topics:

- discrete_variables
- state_variables

**Specification**

**Alias:** none

**Argument(s):** none

Child Keywords:
7.4 variables

### Description

Discrete state variables whose values come from a set of admissible elements. Each variable specified must be of type integer, string, or real.

### 7.4.32.1 integer

- **Keywords Area**
  - variables
  - discrete_state_set
  - integer

Discrete state variables, each defined by a set of permissible integers

### Topics

This keyword is related to the topics:

- discrete_variables
- state_variables

### Specification

**Alias:** none

**Argument(s):** INTEGER

**Default:** no discrete state set integer variables

**Child Keywords:**

---

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<tr>
<td>Optional</td>
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<td></td>
<td>Labels for the variables</td>
</tr>
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</table>

**Description**

Discrete state variables defined by a set of permissible integers.

The details of how to specify this discrete variable are located on the `discrete_variables` page.

See the `state_variables` page for details on the behavior of state variables.

7.4.32.1.1 `elements_per_variable`

- Keywords Area
- variables
- discrete_state_set
- integer
- elements_per_variable

Number of admissible elements for each set variable

**Specification**

**Alias:** num_set_values

**Argument(s):** INTEGERLIST

**Default:** equal distribution

**Description**

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.
The permissible values for each discrete variable

**Specification**

**Alias:** set_values

**Argument(s):** INTEGERLIST

**Description**

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

Whether the set-valued variables are categorical or relaxable

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Description**

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include 'yes', 'no', 'true', and 'false', or any abbreviation in [yYnNtTfF]+

**Examples**

Discrete_design_set variable, 'rotor_blades', can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and 'rotor_blades' can take on any value between 2 and 7, e.g., 3, 6, or 5.5.
discrete_design_set
  integer 1
  elements 2 4 7
  descriptor 'rotor_blades'
categorical 'no'

7.4.32.1.4 initial_state

- Keywords Area
- variables
- discrete_state_set
- integer
- initial_state

Initial values for variables

Specification

Alias: none
Argument(s): INTEGERLIST
Default: middle set value, or rounded down

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the
value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range
are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of
the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any speci-
fied lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic
assumption on the specified intervals or set values.

7.4.32.1.5 descriptors

- Keywords Area
- variables
- discrete_state_set
- integer
- descriptors

Labels for the variables
7.4 variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: dssiv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.32.2 string

• Keywords Area
• variables
• discrete_state_set
• string

String-valued discrete state set variables

Topics

This keyword is related to the topics:

• discrete_variables
• state_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no discrete state set string variables

Child Keywords:

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<tr>
<td>Optional</td>
<td>elements_per_variable</td>
<td></td>
<td>Number of admissible elements for each set variable</td>
</tr>
</tbody>
</table>
### Description

Discrete state variables whose values come from a specified set of admissible strings. The details of how to specify this discrete variable are located on the `discrete_variables` page. See the `state_variables` page for details on the behavior of state variables. Each string element value must be quoted and may contain alphanumeric, dash, underscore, and colon. White space, quote characters, and backslash/metacharacters are not permitted.

#### 7.4.32.2.1 elements_per_variable

- **Keywords Area**
  - variables
  - discrete_state_set
  - string
  - elements_per_variable

Number of admissible elements for each set variable

#### Specification

**Alias:** num_set_values

**Argument(s):** INTEGERLIST

**Default:** equal distribution

#### Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.

#### 7.4.32.2.2 elements

- **Keywords Area**
  - variables
  - discrete_state_set
  - string
  - elements

The permissible values for each discrete variable
7.4 variables

Specification

**Alias:** set_values

**Argument(s):** STRINGLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the **discrete_variables** page.

7.4.32.2.3 initial_state

- **Keywords Area**
- **variables**
- **discrete_state_set**
- **string**
- **initial_state**

Initial values for variables

Specification

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** middle set value, or rounded down

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

**Default Behavior**

Range variables, including **continuous_design**, **continuous_state**, **discrete_design_range**, and **discrete_state_range** are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including **discrete_design_set** and **discrete_state_set** default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.
Keywords Area

7.4.32.2.4 descriptors

- Keywords Area
- variables
- discrete_state_set
- string
- descriptors

Labels for the variables

Specification

Alias: none
Argument(s): STRINGLIST
Default: dsssv_{(i)}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.32.3 real

- Keywords Area
- variables
- discrete_state_set
- real

Discrete state variables, each defined by a set of permissible real numbers

Topics

This keyword is related to the topics:

- discrete_variables
- state_variables

Specification

Alias: none
Argument(s): INTEGER
Default: no discrete state set real variables
Child Keywords:
### 7.4 variables

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</tr>
<tr>
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<td></td>
<td>categorical</td>
<td>Whether the set-valued variables are categorical or relaxable</td>
</tr>
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<tr>
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<td></td>
<td>descriptors</td>
<td>Labels for the variables</td>
</tr>
</tbody>
</table>

#### Description

Discrete state variables defined by a set of permissible real numbers.

The details of how to specify this discrete variable are located on the discrete_variables page.

See the state_variables page for details on the behavior of state variables.

#### 7.4.32.3.1 elements_per_variable

- **Keywords Area**
- variables
- discrete_state_set
- real
- elements_per_variable

Number of admissible elements for each set variable

#### Specification

**Alias:** num_set_values

**Argument(s):** INTEGERLIST

**Default:** equal distribution

#### Description

Discrete set variables (including design, uncertain, and state) take on only a fixed set of values. For each type (integer, string, or real), this keyword specifies how many admissible values are provided for each variable. If not specified, equal apportionment of elements among variables is assumed, and the number of elements must be evenly divisible by the number of variables.
7.4.32.3.2  elements

• Keywords Area
• variables
• discrete_state_set
• real
• elements

The permissible values for each discrete variable

Specification

Alias: set_values
Argument(s): REALLIST

Description

Specify the permissible values for discrete set variables (of type integer, string, or real). See the description on the discrete_variables page.

7.4.32.3.3  categorical

• Keywords Area
• variables
• discrete_state_set
• real
• categorical

Whether the set-valued variables are categorical or relaxable

Specification

Alias: none
Argument(s): STRINGLIST

Description

A list of strings of length equal to the number of set (integer, string, or real) variables indicating whether they are strictly categorical, meaning may only take on values from the provided set, or relaxable, meaning may take on any integer or real value between the lowest and highest specified element. Valid categorical strings include 'yes', 'no', 'true', and 'false', or any abbreviation in [yYnNtTfF]

Examples

Discrete_design_set variable, 'rotor_blades', can take on only integer values, 2, 4, or 7 by default. Since categorical is specified to be false, the integrality can be relaxed and 'rotor_blades' can take on any value between 2 and 7, e.g., 3, 6, or 5.5.
discrete_design_set
integer 1
   elements 2 4 7
descriptor 'rotor_blades'
categorical 'no'

7.4.32.3.4 initial_state

- Keywords Area
- variables
- discrete_state_set
- real
- initial_state

Initial values for variables

Specification

Alias: none

Argument(s): REALLIST

Default: middle set value, or rounded down

Description

Specifies the initial or default value a variable should assume, for example when starting a method or fixing the value of an inactive variable.

Default Behavior

Range variables, including continuous_design, continuous_state, discrete_design_range, and discrete_state_range are initialized to:

- Mean/midpoint (or next smaller integer value) when doubly-bounded
- 0 when unbounded
- 0 when semi-bounded and zero is in the half interval, otherwise to the specified lower or upper bound

Set variables, including discrete_design_set and discrete_state_set default to the middle (or next smaller) value of the ordered set of admissible values.

Uncertain variables default initialize to their mean (or nearest admissible discrete) value, truncating to any specified lower or upper bounds as needed. Epistemic uncertain variable means are calculated making a probabilistic assumption on the specified intervals or set values.

7.4.32.3.5 descriptors

- Keywords Area
- variables
- discrete_state_set
- real
- descriptors

Labels for the variables
Specification

Alias: none
Argument(s): STRINGLIST
Default: dssrv_{i}

Description

The optional variable labels specification descriptors is a list of strings which identify the variables. These are used in console and tabular output.

The default descriptor strings use a variable type-dependent root string plus a numeric identifier.

7.4.33 linear_inequality_constraint_matrix

- Keywords Area
- variables
- linear_inequality_constraint_matrix

Define coefficients of the linear inequality constraints

Topics

This keyword is related to the topics:

- linear_constraints

Specification

Alias: none
Argument(s): REALLIST
Default: no linear inequality constraints

Description

In the inequality case, the constraint matrix $A$ provides coefficients for the variables in the two-sided formulation:

$$a_l \leq Ax \leq a_u$$

Where the bounds are optionally specified by linear_inequality_lower_bounds, and linear_inequality_upper_bounds. The bounds, if not specified, will default to -infinity, and 0, respectively, resulting in one-sided inequalities of the form

$$Ax \leq 0.0$$

The linear_constraints topics page (linked above) outlines a few additional things to consider when using linear constraints.
Examples

In the first example, an optimization involving two variables, \(x_1\) and \(x_2\), is to be performed. These variables must satisfy two constraints:

\[
1.5 \cdot x_1 + 1.0 \cdot x_2 \leq 5.0 \\
x_1 \leq x_2 \implies x_1 - x_2 \leq 0.0
\]

The pair of constraints can be written in matrix form as:

\[
\begin{bmatrix}
1.5 & 1.0 \\
1.0 & -1.0
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2
\end{bmatrix}
\leq
\begin{bmatrix}
5.0 \\
0.0
\end{bmatrix}
\]

The coefficient matrix and right hand side of this matrix inequality are expressed to Dakota in the variables section of the input file:

```
variables
  continuous_design 2
  descriptors 'x1' 'x2'
linear_inequality_constraint_matrix = 1.5 1.0
  1.0 -1.0
linear_inequality_upper_bounds = 5.0
  0.0
```

The second example is more complex in two respects. First, some, but not all, of the constraints are “two sided”, with both lower and upper bounds. Second, not all variables participate in all constraints.

There are four variables, \(x_1\), \(x_2\), \(x_3\), and \(x_4\), and four constraints.

\[
-2.0 \leq 5.0 \cdot x_1 + 2.0 \cdot x_2 \leq 9.0 \\
0.0 \leq x_1 + x_3 \\
-8.0 \leq x_2 + 6.0 \cdot x_4 \leq 8.0 \\
x_1 + x_2 + x_3 \leq 9.0
\]

Or, in matrix form,

\[
\begin{bmatrix}
-2.0 \\
0.0 \\
-8.0 \\
-\infty
\end{bmatrix}
\leq
\begin{bmatrix}
5.0 & 2.0 & 0.0 & 0.0 \\
1.0 & 0.0 & 1.0 & 0.0 \\
0.0 & 1.0 & 0.0 & 6.0 \\
1.0 & 1.0 & 1.0 & 0.0
\end{bmatrix}
\begin{bmatrix}
x_1 \\
x_2 \\
x_3 \\
x_4
\end{bmatrix}
\leq
\begin{bmatrix}
9.0 \\
\infty \\
8.0 \\
9.0
\end{bmatrix}
\]

The Dakota specification for this matrix inequality is:

```
variables
  continuous_design 4
  descriptors 'x1' 'x2' 'x3' 'x4'
linear_inequality_constraint_matrix = 5.0 2.0 0.0 0.0
  1.0 0.0 1.0 0.0 \\
  0.0 1.0 0.0 6.0 \\
  1.0 1.0 1.0 0.0
linear_inequality_lower_bounds = -2.0
  -8.0
linear_inequality_upper_bounds = 9.0
  8.0
```

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7.4.34  linear_inequality_lower_bounds

- Keywords Area
- variables
- linear_inequality_lower_bounds

Define lower bounds for the linear inequality constraint

Topics

This keyword is related to the topics:

- linear_constraints

Specification

Alias: none

Argument(s): REALLIST

Default: vector values = -infinity

Description

In the inequality case, the lower $a_l$ and upper $a_u$ bounds provide constraint limits for the two-sided formulation:

$$a_l \leq Ax \leq a_u$$

Where $A$ is the constrain matrix of variable coefficients.

As with nonlinear inequality constraints (see objective_functions), the default linear inequality constraint bounds are selected so that one-sided inequalities of the form

$$Ax \leq 0$$

result when there are no user bounds specifications (this provides backwards compatibility with previous Dakota versions).

In a user bounds specification, any upper bound values greater than $+\text{bigRealBoundSize}$ (1.e+30, as defined in Minimizer) are treated as $+\infty$ and any lower bound values less than $-\text{bigRealBoundSize}$ are treated as $-\infty$.

This feature is commonly used to drop one of the bounds in order to specify a 1-sided constraint (just as the default lower bounds drop out since $-\text{DBL_MAX} < -\text{bigRealBoundSize}$).

Examples

Examples of specifying linear inequality constraints to Dakota are provided on the linear_inequality_constraint_matrix page.

7.4.35  linear_inequality_upper_bounds

- Keywords Area
- variables
- linear_inequality_upper_bounds

Define upper bounds for the linear inequality constraint
7.4 variables

Topics

This keyword is related to the topics:

- linear_constraints

Specification

Alias: none

Argument(s): REALLIST

Default: vector values = 0.

Description

In the inequality case, the lower $a_l$ and upper $a_u$ bounds provide constraint limits for the two-sided formulation:

$$a_l \leq Ax \leq a_u$$

Where $A$ is the constrain matrix of variable coefficients.

As with nonlinear inequality constraints (see objective_functions), the default linear inequality constraint bounds are selected so that one-sided inequalities of the form

$$Ax \leq 0.0$$

result when there are no user bounds specifications (this provides backwards compatibility with previous Dakota versions).

In a user bounds specification, any upper bound values greater than $+\text{bigRealBoundSize}$ ($1.0e+30$, as defined in Minimizer) are treated as $+\text{infinity}$ and any lower bound values less than $-\text{bigRealBoundSize}$ are treated as $-\text{infinity}$.

This feature is commonly used to drop one of the bounds in order to specify a 1-sided constraint (just as the default lower bounds drop out since $-\text{DBL_MAX} < -\text{bigRealBoundSize}$).

Examples

Examples of specifying linear inequality constraints to Dakota are provided on the linear_inequality_constraint_matrix page.

7.4.36 linear_inequality_scale_types

- Keywords Area
- variables
- linear_inequality_scale_types

How to scale each linear inequality constraint

Topics

This keyword is related to the topics:

- linear_constraints
Specification

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** vector values = "none"

**Description**

Each string in `linear_inequality_scale_types` indicates the scaling type for each linear inequality constraint. They only have effect when the associated method specifies scaling.

The options are:

- ‘value’ - characteristic value. If this is chosen, then `linear_inequality_scales` must be specified; ‘value’ is assumed if scales are given without `scale_types`
- ‘auto’ - automatic scaling.

If a single string is specified it will apply to all linear inequality constraints. Otherwise the number of strings specified should be equal to the number of linear inequalities.

Scaling for linear constraints is applied after any continuous variable scaling.

For example, for variable scaling on continuous design variables $x$:

$$\tilde{x}^j = \frac{x^j - x^j_O}{x^j_M}$$

we have the following system for linear inequality constraints

$$a_L \leq A_i x \leq a_U$$

$$a_L \leq A_i (\text{diag}(x_M)\tilde{x} + x_O) \leq a_U$$

$$a_L - A_i x_O \leq A_i \text{diag}(x_M)\tilde{x} \leq a_U - A_i x_O$$

$$\tilde{a}_L \leq \tilde{A}_i \tilde{x} \leq \tilde{a}_U$$

and user-specified or automatically computed scaling multipliers are applied to this final transformed system, which accounts for continuous design variable scaling. When automatic scaling is in use for linear constraints they are linearly scaled by a computed characteristic value, but not affinely to [0,1].

See the scaling information under specific methods, e.g., `method-#-scaling` for details on how to use this keyword.

### 7.4.37 linear_inequality_scales

- **Keywords Area**
- **variables**
- **linear_inequality_scales**

Characteristic values to scale linear inequalities

**Topics**

This keyword is related to the topics:

- **linear_constraints**
7.4 variables

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** vector values = 1 . (no scaling)

**Description**

Each real value in `linear_inequality_scales` is a nonzero characteristic value to be used in scaling each constraint. They only have effect when the associated method specifies scaling.

This keyword is required for `linear_inequality_scale_types` of ‘value’ and optional for ‘auto’.

If a single real value is specified it will apply to all linear inequality constraints. Otherwise the number of values should be equal to the number of linear inequalities.

Scaling for linear constraints is applied after any continuous variable scaling.

For example, for variable scaling on continuous design variables \( x \):

\[
\tilde{x}^j = \frac{x^j - x^j_O}{x^j_M - x^j_O}
\]

we have the following system for linear inequality constraints

\[
\begin{align*}
a_L & \leq A_i x \leq a_U \\
a_L & \leq A_i (\text{diag}(x_M)\tilde{x} + x_O) \leq a_U \\
a_L - A_i x_O & \leq A_i \text{diag}(x_M)\tilde{x} \leq a_U - A_i x_O \\
\tilde{a}_L & \leq \tilde{A}_i \tilde{x} \leq \tilde{a}_U
\end{align*}
\]

and user-specified or automatically computed scaling multipliers are applied to this final transformed system, which accounts for continuous design variable scaling. When automatic scaling is in use for linear constraints they are linearly scaled by a computed characteristic value, but not affinely to \([0,1]\).

See the scaling information under specific methods, e.g., `method-*=scaling` for details on how to use this keyword.

7.4.38 linear_equality_constraint_matrix

- **Keywords Area**
- **variables**
  - `linear_equality_constraint_matrix`

Define coefficients of the linear equalities

**Topics**

This keyword is related to the topics:

- `linear_constraints`

**Specification**

**Alias:** none

**Argument(s):** REALLIST

**Default:** no linear equality constraints
Description

In the equality case, the constraint matrix $A$ provides coefficients for the variables on the left hand side of:

$$Ax = a_t$$

The linear_constraints topics page (linked above) outlines a few additional things to consider when using linear constraints.

Examples

An optimization involving three variables, $x_1$, $x_2$, and $x_3$, is to be performed. These variables must satisfy a pair of linear equality constraints:

1. $1.5 \cdot x_1 + 1.0 \cdot x_2 = 5.0$
2. $3.0 \cdot x_1 - 4.0 \cdot x_3 = 0.0$

The pair of constraints can be written in matrix form as:

$$\begin{bmatrix} 1.5 & 1.0 & 0.0 \\ 3.0 & 0.0 & -4.0 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 5.0 \\ 0.0 \end{bmatrix}$$

The coefficient matrix and right hand side are expressed to Dakota in the variables section of the input file:

```plaintext
variables
  continuous_design 2
  descriptors 'x1' 'x2'
linear_equality_constraint_matrix = 1.5 1.0 0.0
                                3.0 0.0 -4.0
linear_equality_targets = 5.0 0.0
```

For related examples, see the linear_inequality_constraint_matrix keyword page.

7.4.39 linear_equality_targets

- Keywords Area
- variables
  - linear_equality_targets

Define target values for the linear equality constraints

Topics

This keyword is related to the topics:

- linear_constraints
7.4 variables

Specification

Alias: none
Argument(s): REALIST
Default: vector values = 0.

Description

In the equality case, the targets \( a_t \) provide the equality constraint right hand sides:

\[ Ax = a_t \]

If this is not specified, the defaults for the equality constraint targets enforce a value of 0. for each constraint:

\[ Ax = 0. \]

Examples

Examples of specifying linear equality constraints to Dakota are provided on the linear_equality_constraint_matrix page.

7.4.40 linear_equality_scale_types

- Keywords Area
- variables
- linear_equality_scale_types

How to scale each linear equality constraint

Topics

This keyword is related to the topics:

- linear_constraints

Specification

Alias: none
Argument(s): STRINGLIST
Default: vector values = "none"

Description

Each string in linear_equality_scale_types indicates the scaling type for each linear equality constraint. They only have effect when the associated method specifies scaling.

The options are:

- ‘value’ - characteristic value. If this is chosen, then linear_equality_scales must be specified; ‘value’ is assumed if scales are given without a scale_types
• ‘auto’ - automatic scaling.

If a single string is specified it will apply to all linear equality constraints. Otherwise the number of strings specified should be equal to the number of linear equalities.

Scaling for linear constraints is applied after any continuous variable scaling.

For example, for variable scaling on continuous design variables x:

\[ \tilde{x}_j = \frac{x_j - x_{jO}}{x_{jM}} \]

we have the following system for linear equality constraints

\[ a_L \leq A_i x \leq a_U \]
\[ a_L \leq A_i (\text{diag}(x_M)\tilde{x} + x_{jO}) \leq a_U \]
\[ a_L - A_i x_{jO} \leq A_i \text{diag}(x_M)\tilde{x} \leq a_U - A_i x_{jO} \]
\[ \tilde{a}_L \leq \tilde{A}_i \tilde{x} \leq \tilde{a}_U \]

and user-specified or automatically computed scaling multipliers are applied to this final transformed system, which accounts for continuous design variable scaling. When automatic scaling is in use for linear constraints they are linearly scaled by a computed characteristic value, but not affinely to [0,1].

See the scaling information under specific methods, e.g., method- scaling for details on how to use this keyword.

### 7.4.41 linear_equality_scales

- Keywords Area
- variables
- linear_equality_scales

Characteristic values to scale linear equalities

**Topics**

This keyword is related to the topics:

- linear_constraints

**Specification**

**Alias**: none

**Argument(s)**: REALLIST

**Default**: vector values = 1. (no scaling)

**Description**

Each real value in linear_equality_scales is a nonzero characteristic value to be used in scaling each constraint. They only have effect when the associated method specifies scaling.

This keyword is required for linear_equality_scale_types of ‘value’ and optional for ‘auto’. 
If a single real value is specified it will apply to all linear equality constraints. Otherwise the number of values should be equal to the number of linear equalities.

Scaling for linear constraints is applied after any continuous variable scaling.

For example, for variable scaling on continuous design variables $x$:

$$
\tilde{x}^j = \frac{x^j - x^j_O}{x^j_M - x^j_O}
$$

we have the following system for linear inequality constraints

$$
a_L \leq A_i x \leq a_U
$$

$$
a_L \leq A_i (\text{diag}(x_M) \tilde{x} + x_O) \leq a_U
$$

$$
a_L - A_i x_O \leq A_i \text{diag}(x_M) \tilde{x} \leq a_U - A_i x_O
$$

$$
\tilde{a}_L \leq \tilde{A}_i \tilde{x} \leq \tilde{a}_U
$$

and user-specified or automatically computed scaling multipliers are applied to this final transformed system, which accounts for continuous design variable scaling. When automatic scaling is in use for linear constraints they are linearly scaled by a computed characteristic value, but not affinely to $[0,1]$.

See the scaling information under specific methods, e.g., method-*-scaling for details on how to use this keyword.

7.5 interface

- Keywords Area

**interface**

Specifies how function evaluations will be performed in order to map the variables into the responses.

Topics

This keyword is related to the topics:

- block

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>id_interface</td>
<td></td>
<td>Name the interface block; helpful when there are multiple</td>
</tr>
<tr>
<td>Optional</td>
<td>analysis_drivers</td>
<td>Define how Dakota should run a function evaluation</td>
<td></td>
</tr>
<tr>
<td>----------</td>
<td>------------------</td>
<td>---------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>algebraic_-</td>
<td>mappings</td>
<td>Use AMPL to define algebraic input-output mappings</td>
</tr>
<tr>
<td>Optional</td>
<td>failure_capture</td>
<td>Determine how Dakota responds to analysis driver failure</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>deactivate</td>
<td>Deactivate Dakota interface features for simplicity or efficiency</td>
<td></td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>batch</td>
<td>Perform evaluations in batches</td>
<td></td>
</tr>
<tr>
<td></td>
<td>asynchronous</td>
<td>Specify local evaluation or analysis concurrency</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>evaluation_servers</td>
<td>Specify the number of evaluation servers when Dakota is run in parallel</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>evaluation_-</td>
<td>scheduling</td>
<td>Specify the scheduling of concurrent evaluations when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>processors_per_-</td>
<td>evaluation</td>
<td>Specify the number of processors per evaluation server when Dakota is run in parallel</td>
</tr>
<tr>
<td>Optional</td>
<td>analysis_servers</td>
<td>Specify the number of analysis servers when Dakota is run in parallel</td>
<td></td>
</tr>
</tbody>
</table>
Description

The interface section in a Dakota input file specifies how function evaluations will be performed in order to map the variables into the responses. The term "interface" refers to the bridge between Dakota and the underlying simulation code.

In this context, a "function evaluation" is the series of operations that takes the variables and computes the responses. This can be comprised of one or many codes, scripts, and glue, which are generically referred to as "analysis drivers" (and optional input/output filters). The mapping actions of analysis_drivers may be combined with explicit algebraic_mappings.

Parallelism Options

- The asynchronous keyword enables concurrent local function evaluations or analyses via operating system process management. Its child keywords allow tailoring the evaluation and analysis concurrency.

- The evaluation servers, scheduling mode (master, peer static or dynamic), and processor keywords allow a user to override Dakota’s default evaluation configuration when running in parallel (MPI) mode.

- The analysis servers and scheduling mode (master, peer static or dynamic) keywords allow a user to override Dakota’s default analysis configuration when running in parallel (MPI) mode.

Note: see direct for the specific processors_per_analysis specification supported for direct interfaces.


Theory

Function evaluations are performed using either interfaces to simulation codes, algebraic mappings, or a combination of the two.

When employing mappings with simulation codes, the interface invokes the simulation using either forks, direct function invocations, or computational grid invocations.

- In the fork case, Dakota will treat the simulation as a black-box and communication between Dakota and the simulation occurs through parameter and result files. This is the most common case.

- In the direct function case, the simulation is internal to Dakota and communication occurs through the function parameter list. The direct case can involve linked simulation codes or test functions which are compiled into the Dakota executable. The test functions allow for rapid testing of algorithms without process creation overhead or engineering simulation expense.

- The grid case is deprecated, but was an experiment in interfacing Dakota to distributed computing engines.

When employing algebraic mappings, the AMPL solver library[29] is used to evaluate a directed acyclic graph (DAG) specification from a separate stub.nl file. Separate stub.col and stub.row files are also required to declare the string identifiers of the subset of inputs and outputs, respectively, that will be used in the algebraic mappings.
7.5.1 id_interface

- Keywords Area
- interface
- id_interface

Name the interface block; helpful when there are multiple

Topics

This keyword is related to the topics:

- block_identifier

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** use of last interface parsed

Description

The optional id_interface keyword accepts a string that uniquely identifies this interface block. A model can then use this interface by specifying the same string in its interface_pointer specification.

**Default Behavior**

If the id_interface specification is omitted, a particular interface specification will be used by a model only if that model does not include an interface_pointer and the interface block was the last (or only) one parsed.

**Usage Tips**

- It is a best practice to always use explicit interface IDs and pointers to avoid confusion.
- If only one interface block exists, then id_interface can be safely omitted from the interface block (and interface_pointer omitted from the model specification(s)), since there is no ambiguity.

**Examples**

For example, a model specification including

```model
  interface_pointer = 'I1'
```

will link to an interface with

```id_interface = 'I1'
```

7.5.2 analysis_drivers

- Keywords Area
- interface
- analysis_drivers

Define how Dakota should run a function evaluation
### Specification

**Alias:** none  
**Argument(s):** STRINGLIST  

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>input_filter</td>
<td>Run a pre-processing script before the analysis drivers</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>output_filter</td>
<td>Run a post-processing script after the analysis drivers (Not recommended) Launch analysis drivers with a system call</td>
</tr>
<tr>
<td>Required (Group 1)</td>
<td>Interface Type</td>
<td>system</td>
<td>Launch analysis drivers using fork command</td>
</tr>
<tr>
<td>Required (Group 1)</td>
<td></td>
<td>fork</td>
<td>Launch analysis drivers using fork command</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>direct</td>
<td>Run analysis drivers that are linked-to or compiled-with Dakota</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>matlab</td>
<td>Run Matlab through a direct interface - requires special Dakota build</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>pybind11</td>
<td>Interface directly to a Python simulation via pybind11 - requires special Dakota build</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>python</td>
<td>Run Python through a direct interface - requires special Dakota build</td>
</tr>
</tbody>
</table>
The `analysis_drivers` keyword provides the names of one or more executable analysis programs or scripts, a.k.a. "drivers" which comprise a function evaluation. The optional and required sub-keywords specify how Dakota will manage directories and files, and run the driver(s).

**Types of Interfaces**

Dakota has two recommended ways of running analysis drivers:

- as an external processes (fork), or
- using internal code to couple to the analysis driver (direct)

Other options are available for advanced users, and are not as well documented, supported, or tested:

- external processes (system)
- internal coupling (python, matlab, scilab, grid)

**Use Cases**

The internally coupled codes have few options because many of the details are already handled with the coupling. Their behavior is described in the `direct` keyword.

For external processes using the `fork` keyword,

A function evaluation may comprise:

1. **A single analysis driver**: Function evaluation, including all pre- and post-processing is contained entirely within a single script/executable.
2. **A single analysis driver with filters**: Function evaluation is explicitly split into pre-processing (performed by the input filter), analysis, and post-processing (by the output filter).
3. **A single analysis driver with environment variables**: Function evaluation is contained within one analysis driver, but it requires environment variables to be set before running.
4. **Multiple analysis drivers**: Drivers are run sequentially or concurrently (See the `asynchronous` keyword) and can have any of the above options as well.

For fork and system interfaces, the `analysis_driver` list contains the names of one or more executable programs or scripts taking parameters files as input and producing results files as output. The first field in each analysis driver string must be an executable program or script for Dakota to spawn to perform the function evaluation. Drivers support:

- One set of nested quotes, for arguments with spaces
- Dakota will define special environment variables `DAKOTA_PARAMETERS_FILE` and `DAKOTA_RESULT S_FILE` which can be used in the driver script.
• Dakota will replace the tokens {PARAMETERS} and {RESULTS} in an analysis driver string with the names of the parameters and results files for that analysis/evaluation. Along with the verbatim keyword, which prevents Dakota from appending the names of the parameters and results files as command line arguments, this feature provides users with greater control over how their analysis drivers are invoked by Dakota.

• Variable definitions preceding the executable program or script, such as 'MY_VAR=2 run_analysis.sh' are no longer supported.

For details and examples see the Simulation Interface Components section of the Interfaces chapter of the User’s Manual; for details on the filters and environment variables, see the subsection on Syntax for Filter and Driver Strings.

Examples

Examples:

1. analysis_drivers = 'run_simulation_part1.sh' 'run_simulation_part2.sh'

2. analysis_driver = 'run_simulation.sh -option "option 1"'

3. analysis_driver = 'simulation.exe -option value -dakota_params $DAKOTA_PARAMETERS_FILE -input sim.in -dakota_results_file $DAKOTA_RESULTS_FILE'

FAQ

Where will Dakota look for the analysis_driver? Dakota will locate analysis_driver programs first in (or relative to) the present working directory (".", the interface-analysis_drivers-fork-work_directory if used, otherwise the directory in which Dakota is started), then the directory from which Dakota is started, then using the system $PATH environment variable (Path% on Windows).

Where should the driver be located? When the driver is a script it is most commonly placed in the same directory as the Dakota input file. When using a work_directory, Dakota will also look for drivers in the specified working directory, so link_files or copy_files may specify the driver to get copied or linked into the work directory. When executable programs are used as drivers, they are often elsewhere on the filesystem. These can be specified using absolute paths, or by prepending the PATH environment variable so Dakota finds them.

What if Dakota fails to run my analysis_driver? Prepend the absolute location of the driver to the PATH environment variable before running Dakota, or specify an absolute path to the driver in the Dakota input file.

7.5.2.1 input_filter

• Keywords Area

• interface

• analysis_drivers

• input_filter

Run a pre-processing script before the analysis drivers

Specification

Alias: none

Argument(s): STRING

Default: no input filter
Description

The optional \texttt{input\_filter} and \texttt{output\_filter} specifications provide the names of separate pre- and post-processing programs or scripts which assist in mapping Dakota parameters files into analysis input files and mapping analysis output files into Dakota results files, respectively.

If there is only a single analysis driver, then it is usually most convenient to combine pre- and post-processing requirements into a single analysis driver script and omit the separate input and output filters. However, in the case of multiple analysis drivers, the input and output filters provide a convenient location for non-repeated pre- and post-processing requirements. That is, input and output filters are only executed once per function evaluation, regardless of the number of analysis drivers, which makes them convenient locations for data processing operations that are shared among the analysis drivers.

The \texttt{verbatim} keyword applies to input and output filters as well as analysis drivers, and Dakota also will substitute the names of the parameters and results files for the tokens \texttt{(PARAMETERS)} and \texttt{(RESULTS)} in input and output filter strings, as explained in the documentation for \texttt{analysis\_drivers}.

7.5.2.2 \ texttt{output\_filter}

- \texttt{Keywords Area}
- \texttt{interface}
- \texttt{analysis\_drivers}
- \texttt{output\_filter}

Run a post-processing script after the analysis drivers

Specification

\textbf{Alias}: none

\textbf{Argument(s)}: STRING

\textbf{Default}: no output filter

Description

The optional \texttt{input\_filter} and \texttt{output\_filter} specifications provide the names of separate pre- and post-processing programs or scripts which assist in mapping Dakota parameters files into analysis input files and mapping analysis output files into Dakota results files, respectively.

If there is only a single analysis driver, then it is usually most convenient to combine pre- and post-processing requirements into a single analysis driver script and omit the separate input and output filters. However, in the case of multiple analysis drivers, the input and output filters provide a convenient location for non-repeated pre- and post-processing requirements. That is, input and output filters are only executed once per function evaluation, regardless of the number of analysis drivers, which makes them convenient locations for data processing operations that are shared among the analysis drivers.

The \texttt{verbatim} keyword applies to input and output filters as well as analysis drivers, and Dakota also will substitute the names of the parameters and results files for the tokens \texttt{(PARAMETERS)} and \texttt{(RESULTS)} in input and output filter strings, as explained in the documentation for \texttt{analysis\_drivers}.

7.5.2.3 \texttt{system}

- \texttt{Keywords Area}
- \texttt{interface}
(Not recommended) Launch analysis drivers with a system call

### Specification

**Alias:** none  
**Argument(s):** none

#### Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>parameters_file</td>
<td>parameters_file</td>
<td>Specify the name of the parameters file</td>
</tr>
<tr>
<td>Optional</td>
<td>results_file</td>
<td>results_file</td>
<td>Specify the name of the results file</td>
</tr>
<tr>
<td>Optional</td>
<td>file_tag</td>
<td>file_tag</td>
<td>Tag each parameters &amp; results file name with the function evaluation number</td>
</tr>
<tr>
<td>Optional</td>
<td>file_save</td>
<td>file_save</td>
<td>Keep the parameters &amp; results files after the analysis driver completes</td>
</tr>
<tr>
<td>Optional</td>
<td>labeled</td>
<td>labeled</td>
<td>Requires correct function value labels in results file</td>
</tr>
<tr>
<td>Optional</td>
<td>aprepro</td>
<td>aprepro</td>
<td>Write parameters files in APREPRO syntax</td>
</tr>
<tr>
<td>Optional</td>
<td>work_directory</td>
<td>work_directory</td>
<td>Perform each function evaluation in a separate working directory</td>
</tr>
<tr>
<td>Optional</td>
<td>allow_existing_results</td>
<td>allow_existing_results</td>
<td>Change how Dakota deals with existing results files</td>
</tr>
<tr>
<td>Optional</td>
<td>verbatim</td>
<td>verbatim</td>
<td>Specify the command Dakota uses to launch analysis driver(s) and filters</td>
</tr>
</tbody>
</table>

### Description

The system call interface is included in Dakota for portability and backward compatibility. Users are strongly encouraged to use the **fork** interface if possible, reverting to system only when necessary. To enable the system...
call interface, replace the fork keyword with system. All other keywords have identical meanings to those for the fork interface.

See Also

These keywords may also be of interest:

- fork

7.5.2.3.1 parameters_file

- Keywords Area
- interface
- analysis_drivers
- system
- parameters_file

Specify the name of the parameters file

Specification

Alias: none
Argument(s): STRING
Default: Unix temp files

Description

The parameters file is used by Dakota to pass the parameter values to the analysis driver. The name of the file can be optionally specified using the parameters_file keyword.

If this is not specified, the default data transfer files are temporary files with system-generated names (e.g., /tmp/dakota_params_aaaa0886).

7.5.2.3.2 results_file

- Keywords Area
- interface
- analysis_drivers
- system
- results_file

Specify the name of the results file

Specification

Alias: none
Argument(s): STRING
Default: Unix temp files
Description

The results file must be written by the analysis driver. It is read by Dakota to determine the response values for each function evaluation.

The name of the file can be optionally specified using the results_file keyword.

If this is not specified, the default data transfer files are temporary files with system-generated names (e.g., /tmp/dakota_results_aaaa0886).

7.5.2.3.3 file_tag

- Keywords Area
- interface
- analysis_drivers
- system
- file_tag

Tag each parameters & results file name with the function evaluation number

Specification

Alias: none

Argument(s): none

Default: no tagging

Description

If this keyword is used, Dakota will append a period and the function evaluation number to the names of the parameter and results files.

Default Behavior

If this keyword is omitted, the default is no file tagging.

Usage Tips

- File tagging is most useful when multiple function evaluations are running simultaneously using files in a shared disk space. The analysis driver will be able to infer the function evaluation number from the file names.

- Note that when the file_save keyword is used, Dakota automatically renames parameters and results files, giving them tags after execution of the analysis driver if they otherwise would be overwritten by the next evaluation.

Examples

If the following is included in the interface section of the Dakota input:

```
parameters_file = params.in
results_file = results.out
file_tag
```

Then for the 3rd evaluation, Dakota will write params.in.3, and will expect results.out.3 to be written by the analysis driver.
7.5.2.3.4 file_save

- Keywords Area
- interface
- analysis_drivers
- system
- file_save

Keep the parameters & results files after the analysis driver completes

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** file cleanup

**Description**

If file_save is used, Dakota will not delete the parameters and results files after the function evaluation is completed.

The default behavior is NOT to save these files.

If file_tag is not specified and the saved files would be overwritten by a future evaluation, Dakota renames them after the analysis driver has run by tagging them with the evaluation number.

File saving is most useful when debugging the data communication between Dakota and the simulation.

7.5.2.3.5 labeled

- Keywords Area
- interface
- analysis_drivers
- system
- labeled

Requires correct function value labels in results file

**Topics**

This keyword is related to the topics:

- file_formats

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** Function value labels optional
Description

The labeled keyword directs Dakota to enforce a stricter results file format and enables more detailed error reporting.

When the labeled keyword is used, function values in results files must be accompanied by their corresponding descriptors. If the user did not supply response descriptors in her Dakota input file, then Dakota auto-generated descriptors are expected.

Distinct error messages are emitted for function values that are out-of-order, repeated, or missing. Labels that appear without a function value and unexpected data are also reported as errors. Dakota attempts to report all errors in a results file, not just the first it encounters. After reporting results file errors, Dakota aborts.

Labels for analytic gradients and Hessians currently are not supported.

Although the labeled keyword is optional, its use is recommended to help catch and identify problems with results files. The User’s Manual contains further information about the results file format.

Default Behavior

By default, Dakota does not require labels for function values, and ignores them if they are present.

7.5.2.3.6 aprepro

- Keywords Area
- interface
- analysis_drivers
- system
- aprepro

Write parameters files in APREPRO syntax

Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: dprepro

Argument(s): none

Default: standard parameters file format

Description

The format of data in the parameters files can be modified for direct usage with the APREPRO pre-processing tool using the aprepro specification.

Without this keyword, the parameters file are written in DPrePro format. DPrePro is a utility included with Dakota, described in the Users Manual[5].

7.5.2.3.7 work_directory

- Keywords Area
Perform each function evaluation in a separate working directory

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** no work directory  
**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>named</td>
<td>named</td>
<td>The base name of the work directory created by Dakota</td>
</tr>
<tr>
<td>Optional</td>
<td>directory_tag</td>
<td>directory_tag</td>
<td>Tag each work directory with the function evaluation number</td>
</tr>
<tr>
<td>Optional</td>
<td>directory_save</td>
<td>directory_save</td>
<td>Preserve the work directory after function evaluation completion</td>
</tr>
<tr>
<td>Optional</td>
<td>link_files</td>
<td>link_files</td>
<td>Paths to be linked into each working directory</td>
</tr>
<tr>
<td>Optional</td>
<td>copy_files</td>
<td>copy_files</td>
<td>Files and directories to be copied into each working directory</td>
</tr>
<tr>
<td>Optional</td>
<td>replace</td>
<td>replace</td>
<td>Overwrite existing files within a work directory</td>
</tr>
</tbody>
</table>

**Description**

When performing concurrent evaluations, it is typically necessary to cloister simulation input and output files in separate directories to avoid conflicts. When the `work_directory` feature is enabled, Dakota will create a directory for each evaluation, with optional tagging (`directory_tag`) and saving (`directory_save`), as with files, and execute the analysis driver from that working directory.

The directory may be named with a string, or left anonymous to use an automatically-generated directory in the system's temporary file space, e.g., `/tmp/dakota_work_c93vb71z/`. The optional `link_files` and `copy_files` keywords specify files or directories which should appear in each working directory.
When using work_directory, the analysis_drivers may be given by an absolute path, located in (or relative to) the startup directory alongside the Dakota input file, in the list of template files linked or copied, or on the $PATH (Path% on Windows).

7.5.2.3.7.1 named

- Keywords Area
- interface
- analysis_drivers
- system
- work_directory
- named

The base name of the work directory created by Dakota

Specification

**Alias:** none

**Argument(s):** STRING

**Default:** dakota_work_xxxxxxx

Description

The named keyword is followed by a string, indicating the name of the work directory created by Dakota. If relative, the work directory will be created relative to the directory from which Dakota is invoked.

If named is not used, the default work directory is a temporary directory with a system-generated name (e.g., /tmp/dakota_work_c93vb71z/).

See Also

These keywords may also be of interest:

- directory_tag
- directory_save

7.5.2.3.7.2 directory_tag

- Keywords Area
- interface
- analysis_drivers
- system
- work_directory
- directory_tag

Tag each work directory with the function evaluation number
Specification

Alias: dir_tag
Argument(s): none
Default: no work directory tagging

Description

If this keyword is used, Dakota will append a period and the function evaluation number to the work directory names. If this keyword is omitted, the default is no tagging, and the same work directory will be used for ALL function evaluations. Tagging is most useful when multiple function evaluations are running simultaneously.

7.5.2.3.7.3 directory_save

Preserve the work directory after function evaluation completion

Specification

Alias: dir_save
Argument(s): none
Default: remove work directory

Description

By default, when a working directory is created by Dakota using the work_directory keyword, it is deleted after the evaluation is completed. The directory_save keyword will cause Dakota to leave (not delete) the directory.

7.5.2.3.7.4 link_files

Paths to be linked into each working directory
7.5 interface

Specification

Alias: none
Argument(s): STRINGLIST
Default: no linked files

Description

Specifies the paths (files or directories) that will be symbolically linked from each working directory. Wildcards using * and ? are permitted. Linking is space-saving and useful for files not modified during the function evaluation. However, not all filesystems support linking, for example, support on Windows varies.

Examples

Specifying

```
link_files = 'siminput*.in' '/path/to/simdir1' 'simdir2/*'
```

will create copies

```
workdir/siminput*.in  # links to each of rundir / siminput*.in
workdir/simdir1/     # whole directory simdir1 linked
workdir/*            # each entry in directory simdir2 linked
```

7.5.2.3.7.5 copy_files

- Keywords Area
- interface
- analysis_drivers
- system
- work_directory
- copy_files

Files and directories to be copied into each working directory

Specification

Alias: none
Argument(s): STRINGLIST
Default: no copied files

Description

Specifies the files or directories that will be recursively copied into each working directory. Wildcards using * and ? are permitted.
Examples

Specifying

```plaintext
copy_files = 'siminput*.in' '/path/to/simdir1' 'simdir2/*'
```

will create copies

```plaintext
workdir/siminput*.in  # files rundir/siminput*.in copied
workdir/simdir1/      # whole directory simdir1 recursively copied
workdir/*             # contents of directory simdir2 recursively copied
```

where rundir is the directory in which Dakota was started.

7.5.2.3.7.6 replace

- Keywords Area
- interface
- analysis_drivers
- system
- work_directory
- replace

Overwrite existing files within a work directory

Specification

Alias: none
Argument(s): none
Default: do not overwrite files

Description

By default, Dakota will not overwrite any existing files in a work directory. The replace keyword changes this behavior to force overwriting.

7.5.2.3.8 allow_existing_results

- Keywords Area
- interface
- analysis_drivers
- system
- allow_existing_results

Change how Dakota deals with existing results files

Specification

Alias: none
Argument(s): none
Default: results files removed before each evaluation
Description

By default Dakota will remove existing results files before invoking the analysis_driver to avoid problems created by stale files in the current directory. To override this behavior and not delete existing files, specify allow_existing_results.

7.5.2.3.9 verbatim

- Keywords Area
- interface
- analysis_drivers
- system
- verbatim

Specify the command Dakota uses to launch analysis driver(s) and filters

Specification

Alias: none

Argument(s): none

Default: driver/filter invocation syntax augmented with file names

Description

The typical commands that Dakota uses to launch analysis drivers are:

```
> analysis_driver parameters_file_name results_file_name
```

Dakota will automatically arrange the executables and file names. If the analysis driver requires a different syntax, the entire command can be specified as the analysis driver and the verbatim keyword will tell Dakota to use this as the command.

Note, this will not allow the use of file_tag, because the exact command must be specified.
For additional information on invocation syntax, see the Interfaces chapter of the Users Manual[5].

Examples

In the following example, the analysis_driver command is run without any edits from Dakota.

```
interface
  analysis_driver = "matlab -nodesktop -nojvm -r 'MatlabDriver_hardcoded_filenames; exit' "
  fork
    parameters_file 'params.in'
    results_file 'results.out'
  verbatim # this tells Dakota to fork the command exactly as written, instead of appending I/O filenames
```

The -r flag identifies the commands that will be run by matlab. The Matlab script has the parameters_file and results_file names hardcoded, so no additional arguments are required.
7.5.2.4 fork

- Keywords Area
- interface
- analysis_drivers
- fork

Launch analysis drivers using fork command

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>parameters_file</td>
<td></td>
<td>Specify the name of the parameters file</td>
</tr>
<tr>
<td>Optional</td>
<td>results_file</td>
<td></td>
<td>Specify the name of the results file</td>
</tr>
<tr>
<td>Optional</td>
<td>file_tag</td>
<td></td>
<td>Tag each parameters &amp; results file name with the function evaluation number</td>
</tr>
<tr>
<td>Optional</td>
<td>file_save</td>
<td></td>
<td>Keep the parameters &amp; results files after the analysis driver completes</td>
</tr>
<tr>
<td>Optional</td>
<td>labeled</td>
<td></td>
<td>Requires correct function value labels in results file</td>
</tr>
<tr>
<td>Optional</td>
<td>aprepro</td>
<td></td>
<td>Write parameters files in APREPRO syntax</td>
</tr>
<tr>
<td>Optional</td>
<td>work_directory</td>
<td></td>
<td>Perform each function evaluation in a separate working directory</td>
</tr>
<tr>
<td>Optional</td>
<td>allow_existing_results</td>
<td></td>
<td>Change how Dakota deals with existing results files</td>
</tr>
</tbody>
</table>
Optional verbatim

Specify the command Dakota uses to launch analysis driver(s) and filters

Description

The `fork` interface is the most common means by which Dakota launches a separate application analysis process. The `fork` interface is recommended over `system` for most analysis drivers that are external to Dakota, i.e., any driver not linked in via the `direct` interface.

As explained in the Users Manual, the parameters and results file names are passed on the command line to the analysis driver(s). If input/output filters are specified, they will be run before/after the analysis drivers. The `verbatim` keyword is used to modify the default driver/filter commands.

For additional information on invocation syntax, see the Interfaces chapter of the Users Manual[5].

Examples

Spawn (fork) an external executable/script called 'rosenbrock' which reads variables from params.in and writes responses to results.out. Preserve the analysis files for each function evaluation with tag and save.

```plaintext
interface
  analysis_drivers = 'rosenbrock'
  fork
  parameters_file = 'params.in'
  results_file   = 'results.out'
  file_tag
  file_save
```

7.5.2.4.1 parameters_file

- Keywords Area
- interface
- analysis_drivers
- fork
- parameters_file

Specify the name of the parameters file

Specification

Alias: none

Argument(s): STRING

Default: Unix temp files

Description

The parameters file is used by Dakota to pass the parameter values to the analysis driver. The name of the file can be optionally specified using the `parameters_file` keyword.

If this is not specified, the default data transfer files are temporary files with system-generated names (e.g., `/tmp/dakota_params_aaaa0886`).
7.5.2.4.2 results_file

- Keywords Area
- interface
- analysis_drivers
- fork
- results_file

Specify the name of the results file

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** Unix temp files

**Description**

The results file must be written by the analysis driver. It is read by Dakota to determine the response values for each function evaluation.

The name of the file can be optionally specified using the `results_file` keyword.

If this is not specified, the default data transfer files are temporary files with system-generated names (e.g., `/tmp/dakota_results_aaaa0886`).

7.5.2.4.3 file_tag

- Keywords Area
- interface
- analysis_drivers
- fork
- file_tag

Tag each parameters & results file name with the function evaluation number

**Specification**

**Alias:** none

**Argument(s):** none

**Default:** no tagging

**Description**

If this keyword is used, Dakota will append a period and the function evaluation number to the names of the parameter and results files.

**Default Behavior** If this keyword is omitted, the default is no file tagging.

**Usage Tips**
• File tagging is most useful when multiple function evaluations are running simultaneously using files in a shared disk space. The analysis driver will be able to infer the function evaluation number from the file names.

• Note that when the file_save keyword is used, Dakota automatically renames parameters and results files, giving them tags after execution of the analysis driver if they otherwise would be overwritten by the next evaluation.

Examples

If the following is included in the interface section of the Dakota input:

```
parameters_file = params.in
results_file = results.out
file_tag
```

Then for the 3rd evaluation, Dakota will write `params.in.3`, and will expect `results.out.3` to be written by the analysis driver.

7.5.2.4.4 file_save

• Keywords Area
• interface
• analysis_drivers
• fork
• file_save

Keep the parameters & results files after the analysis driver completes

Specification

Alias: none

Argument(s): none

Default: file cleanup

Description

If file_save is used, Dakota will not delete the parameters and results files after the function evaluation is completed.

The default behavior is NOT to save these files.

If file_tag is not specified and the saved files would be overwritten by a future evaluation, Dakota renames them after the analysis driver has run by tagging them with the evaluation number.

File saving is most useful when debugging the data communication between Dakota and the simulation.
• labeled

Requires correct function value labels in results file

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: Function value labels optional

Description

The labeled keyword directs Dakota to enforce a stricter results file format and enables more detailed error reporting.

When the labeled keyword is used, function values in results files must be accompanied by their corresponding descriptors. If the user did not supply response descriptors in her Dakota input file, then Dakota auto-generated descriptors are expected.

Distinct error messages are emitted for function values that are out-of-order, repeated, or missing. Labels that appear without a function value and unexpected data are also reported as errors. Dakota attempts to report all errors in a results file, not just the first it encounters. After reporting results file errors, Dakota aborts.

Labels for analytic gradients and Hessians currently are not supported.

Although the labeled keyword is optional, its use is recommended to help catch and identify problems with results files. The User's Manual contains further information about the results file format.

Default Behavior

By default, Dakota does not require labels for function values, and ignores them if they are present.

7.5.2.4.6 aprepro

• Keywords Area

• interface

• analysis_drivers

• fork

• aprepro

Write parameters files in APREPRO syntax

Topics

This keyword is related to the topics:

• file_formats
7.5 interface

Specification

**Alias:** dprepro

**Argument(s):** none

**Default:** standard parameters file format

Description

The format of data in the parameters files can be modified for direct usage with the APREPRO pre-processing tool [78] using the aprepro specification

Without this keyword, the parameters file are written in DPrePro format. DPrePro is a utility included with Dakota, described in the Users Manual[5].

7.5.2.4.7 work_directory

- Keywords Area
  - interface
  - analysis_drivers
  - fork
  - work_directory

Perform each function evaluation in a separate working directory

Specification

**Alias:** none

**Argument(s):** none

**Default:** no work directory

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>named</td>
<td></td>
<td>The base name of the work directory created by Dakota</td>
</tr>
<tr>
<td>Optional</td>
<td>directory_tag</td>
<td></td>
<td>Tag each work directory with the function evaluation number</td>
</tr>
<tr>
<td>Optional</td>
<td>directory_save</td>
<td></td>
<td>Preserve the work directory after function evaluation completion</td>
</tr>
</tbody>
</table>
When performing concurrent evaluations, it is typically necessary to cloister simulation input and output files in separate directories to avoid conflicts. When the work_directory feature is enabled, Dakota will create a directory for each evaluation, with optional tagging (directory_tag) and saving (directory_save), as with files, and execute the analysis driver from that working directory.

The directory may be named with a string, or left anonymous to use an automatically-generated directory in the system’s temporary file space, e.g., /tmp/dakota_work_c93vb71z/. The optional link_files and copy_files keywords specify files or directories which should appear in each working directory.

When using work_directory, the analysis_drivers may be given by an absolute path, located in (or relative to) the startup directory alongside the Dakota input file, in the list of template files linked or copied, or on the $PATH (Path% on Windows).

7.5.2.4.7.1 named

- Keywords Area
- interface
- analysis_drivers
- fork
- work_directory
- named

The base name of the work directory created by Dakota

**Specification**

**Alias:** none

**Argument(s):** STRING

**Default:** dakota_work_xxxxxxxxx

**Description**

The named keyword is followed by a string, indicating the name of the work directory created by Dakota. If relative, the work directory will be created relative to the directory from which Dakota is invoked.

If named is not used, the default work directory is a temporary directory with a system-generated name (e.g., /tmp/dakota_work_c93vb71z/).
See Also

These keywords may also be of interest:

- directory_tag
- directory_save

7.5.2.4.7.2  directory_tag

- Keywords Area
- interface
- analysis_drivers
- fork
- work_directory
- directory_tag

Tag each work directory with the function evaluation number

Specification

**Alias**: dir_tag

**Argument(s)**: none

**Default**: no work directory tagging

**Description**

If this keyword is used, Dakota will append a period and the function evaluation number to the work directory names. If this keyword is omitted, the default is no tagging, and the same work directory will be used for ALL function evaluations. Tagging is most useful when multiple function evaluations are running simultaneously.

7.5.2.4.7.3  directory_save

- Keywords Area
- interface
- analysis_drivers
- fork
- work_directory
- directory_save

Preserve the work directory after function evaluation completion

Specification

**Alias**: dir_save

**Argument(s)**: none

**Default**: remove work directory
By default, when a working directory is created by Dakota using the `work_directory` keyword, it is deleted after the evaluation is completed. The `directory_save` keyword will cause Dakota to leave (not delete) the directory.

7.5.2.4.7.4 `link_files`

- **Keywords Area**
- **interface**
- **analysis_drivers**
- **fork**
- **work_directory**
- **link_files**

Paths to be linked into each working directory

**Specification**

**Alias:** none

**Argument(s):** STRINGLIST

**Default:** no linked files

**Description**

Specifies the paths (files or directories) that will be symbolically linked from each working directory. Wildcards using * and ? are permitted. Linking is space-saving and useful for files not modified during the function evaluation. However, not all filesystems support linking, for example, support on Windows varies.

**Examples**

Specifying

```plaintext
link_files = 'siminput*.in' '/path/to/simdir1' 'simdir2/*'
```

will create copies

```plaintext
workdir/siminput*.in   # links to each of rundir / siminput*.in
workdir/simdir1/       # whole directory simdir1 linked
workdir/*              # each entry in directory simdir2 linked
```

7.5.2.4.7.5 `copy_files`

- **Keywords Area**
- **interface**
- **analysis_drivers**
- **fork**
- **work_directory**
- **copy_files**

Files and directories to be copied into each working directory
7.5 interface

Specification

Alias: none
Argument(s): STRINGLIST
Default: no copied files

Description

Specifies the files or directories that will be recursively copied into each working directory. Wildcards using * and ? are permitted.

Examples

Specifying

```
    copy_files = 'siminput*.in' '/path/to/simdir1' 'simdir2/*'
```

will create copies

```
    workdir/siminput*.in  # files rundir/siminput*.in copied
    workdir/simdir1/      # whole directory simdir1 recursively copied
    workdir/*             # contents of directory simdir2 recursively copied
```

where rundir is the directory in which Dakota was started.

7.5.2.4.7.6 replace

- Keywords Area
- interface
- analysis_drivers
- fork
- work_directory
- replace

Overwrite existing files within a work directory

Specification

Alias: none
Argument(s): none
Default: do not overwrite files

Description

By default, Dakota will not overwrite any existing files in a work directory. The replace keyword changes this behavior to force overwriting.
7.5.2.4.8 allow_existing_results

- Keywords Area
- interface
- analysis_drivers
- fork
- allow_existing_results

Change how Dakota deals with existing results files

Specification

Alias: none
Argument(s): none
Default: results files removed before each evaluation

Description

By default Dakota will remove existing results files before invoking the analysis_driver to avoid problems created by stale files in the current directory. To override this behavior and not delete existing files, specify allow_existing_results.

7.5.2.4.9 verbatim

- Keywords Area
- interface
- analysis_drivers
- fork
- verbatim

Specify the command Dakota uses to launch analysis driver(s) and filters

Specification

Alias: none
Argument(s): none
Default: driver/filter invocation syntax augmented with file names

Description

The typical commands that Dakota uses to launch analysis drivers are:

> analysis_driver parameters_file_name results_file_name

Dakota will automatically arrange the executables and file names.

If the analysis driver requires a different syntax, the entire command can be specified as the analysis driver and the verbatim keyword will tell Dakota to use this as the command.
Note, this will not allow the use of file_tag, because the exact command must be specified.
For additional information on invocation syntax, see the Interfaces chapter of the Users Manual[5].

Examples

In the following example, the analysis_driver command is run without any edits from Dakota.

```
interface
  analysis_driver = "matlab -nodesktop -nojvm -r 'MatlabDriver_hardcoded_filenames; exit' "
  fork
    parameters_file 'params.in'
    results_file 'results.out'
  verbatim # this tells Dakota to fork the command exactly as written, instead of appending I/O filenames
```

The -r flag identifies the commands that will be run by matlab. The Matlab script has the parameters_file and results_file names hardcoded, so no additional arguments are required.

7.5.2.5 direct

• Keywords Area
• interface
• analysis_drivers
• direct

Run analysis drivers that are linked-to or compiled-with Dakota

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>processors_per_analysis</td>
<td></td>
<td>Specify the number of processors per analysis when Dakota is run in parallel</td>
</tr>
</tbody>
</table>

Description

Direct interfaces are used to compile/link simulation programs into Dakota and to invoke Dakota’s built-in algebraic test problems.

Direct simulation interfaces communicate variable and response data in-core instead of through the filesystem. This typically requires modification to simulator programs so that they can be linked into Dakota; however it can be more efficient due to elimination of external processes and auxiliary simulator output, more accurate due to higher numerics, and more flexible in terms of MPI parallelism.

Direct interfaces are also used to invoke internal test functions that perform parameter to response mappings for simple functions as inexpensively as possible. These problems are compiled directly into the Dakota executable as part of the direct function interface class and are used for algorithm testing.
Dakota supports direct interfaces to a few select simulation codes such as Matlab, Python, and Scilab. Another example is ModelCenter, a commercial simulation management framework from Phoenix Integration. To utilize this interface, a user must first define the simulation specifics within a ModelCenter session and then save these definitions to a ModelCenter configuration file. The `analysis_components` specification provides the means to communicate this configuration file to Dakota’s ModelCenter interface.

Examples

The rosenbrock function is available as an executable, which can be launched with `fork`, and is also compiled with Dakota. The internal version can be used with:

```plaintext
interface
  analysis_drivers = 'rosenbrock'
  direct

7.5.2.5.1 processors_per_analysis
```

Specify the number of processors per analysis when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Default: automatic (see discussion)

Description

For direct function interfaces, `processors_per_analysis` can be used to specify multiprocessor analysis partitions. As with the `evaluation_servers`, `analysis_servers`, `evaluation_self_scheduling`, `evaluation_static_scheduling`, `analysis_self_scheduling`, and `analysis_static_scheduling` specifications, `processors_per_analysis` provides a means for the user to override the automatic parallel configuration (refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5]) for the number of processors used for each analysis partition.

Usage Tips

- If both `analysis_servers` and `processors_per_analysis` are specified and they are not in agreement, then `analysis_servers` takes precedence.
7.5 interface

7.5.2.6 matlab

- Keywords Area
- interface
- analysis_drivers
- matlab

Run Matlab through a direct interface - requires special Dakota build

Specification

Alias: none
Argument(s): none

Description

Dakota supports a library-linked interface to Matlab, but it must be explicitly enabled when compiling Dakota from source. Consult the Users Manual[5] for discussion and examples. Contact the Dakota users mailing list for assistance building and using Dakota with these interfaces.

The analysis_drivers specifies a Matlab file which implements the parameter to response mapping.

Examples

See dakota/share/dakota/examples/users/MATLAB/linked

7.5.2.7 pybind11

- Keywords Area
- interface
- analysis_drivers
- pybind11

Interface directly to a Python simulation via pybind11 - requires special Dakota build

Specification

Alias: none
Argument(s): none

Description

Dakota supports a library-linked interface to Python using the pybind11 C++ API, but it must be explicitly enabled when compiling Dakota from source. This is an emerging capability currently intended to work with a top-level Dakota Python module instantiated from Python, together with one or more registered Python callback functions. The name of the callback function used to map parameters to responses is the argument in the analysis_drivers specification. List data structures are passed to the specified Python callback function with responses expected to be populated therein.
Usage Tips  This is a new capability which should be considered preliminary with limitations, e.g., poor handling of the Global Interpreter Lock can produce failures on Mac platforms.

Examples

See dakota_src/src/unit_test/test_dakota.py

7.5.2.8 python

- Keywords Area
- interface
- analysis_drivers
- python

Run Python through a direct interface - requires special Dakota build

Specification

Alias: none
Argument(s): none
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>numpy</td>
<td>Enable the use of numpy in Dakota’s Python interface</td>
</tr>
</tbody>
</table>

Description

Dakota supports a library-linked interface to Python, but it must be explicitly enabled when compiling Dakota from source. Consult the Users Manual[5] for discussion and examples. Contact the Dakota users mailing list for assistance building and using Dakota with these interfaces.

The analysis_drivers specifies a Python module:function which implements the parameter to response mapping. List data structures are the default, but NumPy is also supported, if enabled in the build.

Examples

See dakota/share/dakota/examples/official/drivers/Python/linked

7.5.2.8.1 numpy

- Keywords Area
- interface
- analysis_drivers
- python
- numpy

Enable the use of numpy in Dakota’s Python interface
7.5 interface

Specification

Alias: none
Argument(s): none
Default: Python list dataflow

Description

When the numpy keyword is used, Dakota expects responses in the form of a Python dictionary of numpy arrays. See the example in dakota/share/dakota/examples/official/drivers/Python/linked.

7.5.2.9 scilab

- Keywords Area
- interface
- analysis_drivers
- scilab

Run Scilab through a direct interface - requires special Dakota build

Specification

Alias: none
Argument(s): none

Description

Dakota supports a library-linked interface to Scilab, but it must be explicitly enabled when compiling Dakota from source. Consult the Users Manual[5] for discussion and examples. Contact the Dakota users mailing list for assistance building and using Dakota with these interfaces.

The analysis_drivers specifies a Scilab file which implements the parameter to response mapping.

Examples

See dakota/share/dakota/examples/users/Scilab/linked

7.5.2.10 grid

- Keywords Area
- interface
- analysis_drivers
- grid

Deprecated grid computing interface
Specification

**Alias**: none

**Argument(s)**: none

**Description**

Interface Dakota directly to a grid (distributed) computing engine. This deprecated capability was historically used for interfaces with IDEA and JAVASpaces in the past and was intended as a placeholder for future work with Condor and/or Globus services. It is not currently operational.

7.5.2.11 analysis_components

- **Keywords Area**
- **interface**
- **analysis_drivers**
- **analysis_components**

Provide additional identifiers to analysis drivers.

**Specification**

**Alias**: none

**Argument(s)**: STRINGLIST

**Default**: no additional identifiers

**Description**

The optional analysis_components specification allows the user to provide additional identifiers (e.g., mesh file names) for use by the analysis drivers. This is particularly useful when the same analysis driver is to be reused multiple times for slightly different analyses. The specific content within the strings is open-ended and can involve whatever syntax is convenient for a particular analysis driver. The number of analysis components $n_c$ should be an integer multiple of the number of drivers $n_d$, and the first $n_c/n_d$ component strings will be passed to the first driver, etc.

7.5.3 algebraic_mappings

- **Keywords Area**
- **interface**
- **algebraic_mappings**

Use AMPL to define algebraic input-output mappings

**Specification**

**Alias**: none

**Argument(s)**: STRING

**Default**: no algebraic mappings
Description

Dakota can evaluate algebraic input-output mappings using AMPL [26]. The mappings are expressed in 3 files: stub.nl, stub.col, and stub.row, where stub is a particular root name describing a particular problem. The file names are communicated to Dakota using the algebraic_mappings keyword. It may either specify the full stub.nl filename, or alternatively, just the stub basename.

Dakota then extracts the input and output identifier strings from stub.col and stub.row and employs the AMPL solver library[29] to process the directed acyclic graph (DAG) specification in stub.nl. The variable and objective function names declared within AMPL should be a subset of the variable and response descriptors specified in the variables and responses blocks. Ordering is not important, as Dakota will reorder data as needed.

Examples

An interface employing both algebraic and simulation-based mappings. The results from the individual algebraic and simulation mappings are overlaid based on the variable and response descriptors used by the individual mappings.

```plaintext
interface,
    algebraic_mappings = 'ampl/fma.nl'
for:
    analysis_driver = 'text_book'
    parameters_file = 'tb.in'
    results_file = 'tb.out'
```

7.5.4 failure_capture

- Keywords Area
  - interface
  - failure_capture

Determine how Dakota responds to analysis driver failure

Specification

Alias: none

Argument(s): none

Default: abort

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Failure Mitigation (Group 1)</th>
<th>Dakota Keyword abort</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>(Default) Abort the Dakota job</td>
</tr>
<tr>
<td>retry</td>
<td></td>
<td></td>
<td>Rerun failed analyses</td>
</tr>
<tr>
<td>recover</td>
<td></td>
<td></td>
<td>Substitute dummy values for the responses</td>
</tr>
</tbody>
</table>
Dakota can deal with analysis failure in a few ways. The first step is that Dakota must detect analysis failure. Importantly, Dakota always expects a results file to be written by the analysis driver, even when a failure has occurred. If the file does not exist when the analysis driver exits, a Dakota error results, causing Dakota itself to terminate. The analysis driver communicates an analysis failure to Dakota by writing a results file beginning with the (case-insensitive) word “fail”. Any file contents after “fail” are ignored.

Once Dakota detects analysis failure, the failure can be mitigated in four ways:

- **abort** (the default)
- **retry**
- **recover**
- **continuation**


### 7.5.4.1 abort

- **Keywords Area**
- **interface**
- **failure_capture**
- **abort**

(Default) Abort the Dakota job

**Specification**

**Alias**: none

**Argument(s)**: none

**Description**

Stop the Dakota job, as well as any other running analysis drivers when a failure is communicated.

### 7.5.4.2 retry

- **Keywords Area**
- **interface**
- **failure_capture**
7.5.3 retry

Rerun failed analyses

Specification

Alias: none

Argument(s): INTEGER

Description

Number of times to retry a failing analysis

7.5.4.3 recover

- Keywords Area
- interface
- failure_capture
- recover

Substitute dummy values for the responses

Specification

Alias: none

Argument(s): REALLIST

Description

When a simulation failure is detected, substitute the provided dummy function values in the response. Gradient and Hessian are not supported.

7.5.4.4 continuation

- Keywords Area
- interface
- failure_capture
- continuation

Cause Dakota to step toward the failed "target" simulation from a nearby successful "source"

Specification

Alias: none

Argument(s): none
Description

When `failure_capture continuation` is enabled and an evaluation fails, then Dakota will attempt to march incrementally from a previous good evaluation (the "source") toward the failing one (the "target"). Further details about the algorithm employed by Dakota are supplied in the User's Manual [5].

7.5.5 deactivate

- Keywords Area
- interface
- deactivate

Deactivate Dakota interface features for simplicity or efficiency

Specification

Alias: none
Argument(s): none
Default: Active set vector control, function evaluation cache, and restart file features are active
Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
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<tr>
<td>Optional</td>
<td>active_set_vector</td>
<td>passive_set</td>
<td>Deactivate the Active Set Vector</td>
</tr>
<tr>
<td>Optional</td>
<td>evaluation_cache</td>
<td></td>
<td>Do not retain function evaluation history in memory</td>
</tr>
<tr>
<td>Optional</td>
<td>strict_cache_.equality</td>
<td>strict_cache_.equality</td>
<td>Do not require strict cache equality when finding duplicates</td>
</tr>
<tr>
<td>Optional</td>
<td>restart_file</td>
<td></td>
<td>Deactivate writing to the restart file</td>
</tr>
</tbody>
</table>

Description

The optional deactivate specification block allows a user to deactivate interface features in order to simplify interface development, increase execution speed, and/or reduce memory and disk requirements. Any or all of these features may be specified concurrently.

- Active set vector (ASV) control: deactivate so that Dakota expects the same response data (all functions, gradients, Hessian) back from the simulation on every evaluation, instead of only those components required by the method for this particular function evaluation.

- Function evaluation cache: save memory by not caching the function evaluation history. May result in additional (duplicate) function evaluations.
• Strict cache equality: allow a relaxed tolerance when detecting duplicate function evaluations. Can be useful when importing data or restarting across machines.

• Restart file: improve efficiency and eliminate restart file storage at the risk of not being able to recover a failed or partial Dakota study.

7.5.5.1 active_set_vector

• Keywords Area
  • interface
  • deactivate
  • active_set_vector

Deactivate the Active Set Vector

Specification

Alias: none

Argument(s): none

Description

Allows the user to turn off any variability in ASV values so that active set logic can be omitted in the user’s simulation interface. This option trades some efficiency for simplicity in interface development.

The default behavior is to request the minimum amount of data required by an algorithm at any given time, which implies that the ASV values may vary from one function evaluation to the next. Since the user’s interface must return the data set requested by the ASV values, this interface must contain additional logic to account for any variations in ASV content.

Deactivating this ASV control causes Dakota to always request a “full” data set (the full function, gradient, and Hessian data that is available from the interface as specified in the responses specification) on each function evaluation.

For example, if ASV control has been deactivated and the responses section specifies four response functions, analytic gradients, and no Hessians, then the ASV on every function evaluation will be \{3 3 3 3\}, regardless of what subset of this data is currently needed. While wasteful of computations in many instances, this simplifies the interface and allows the user to return the same data set on every evaluation. Conversely, if ASV control is active (the default behavior), then the ASV requests in this example might vary from \{1 1 1 1\} to \{2 0 0 2\}, etc., according to the specific data needed on a particular function evaluation. This will require the user’s interface to read the ASV requests and perform the appropriate logic in conditionally returning only the data requested.

Usage Tips

• In general, the default ASV behavior is recommended for the sake of computational efficiency, unless interface development time is a critical concern.

• Whether active or inactive, the data returned to Dakota from the user’s interface must match the ASV passed in, or else a response recovery error will result. However, when the ASV control is deactivated, the ASV values are invariant and need not be checked on every evaluation.

• Deactivating the ASV control can have a positive effect on load balancing for parallel Dakota executions. Thus, there is significant overlap in this ASV control option with speculative gradients. There is also overlap with the mode override approach used with certain optimizers to combine individual value, gradient, and Hessian requests.
7.5.5.2 evaluation_cache

- Keywords Area
- interface
- deactivate
- evaluation_cache

Do not retain function evaluation history in memory

Specification

Alias: none

Argument(s): none

Description

Do not retain the complete function evaluation history in memory.

This can be important for reducing memory requirements in large-scale applications (i.e., applications with a large number of variables or response functions) and for eliminating the overhead of searching for duplicates within the function evaluation cache prior to each new function evaluation (e.g., for improving speed in problems with 1000's of inexpensive function evaluations or for eliminating overhead when performing timing studies).

However, the downside is that unnecessary computations may be performed since duplication in function evaluation requests may not be detected. For this reason, this option is not recommended when function evaluations are costly.

Note: duplication detection within Dakota can be deactivated, but duplication detection features within specific optimizers may still be active.

7.5.5.3 strict_cache_equality

- Keywords Area
- interface
- deactivate
- strict_cache_equality

Do not require strict cache equality when finding duplicates

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>cache_tolerance</td>
<td></td>
<td>Specify tolerance when identifying duplicate function evaluations</td>
</tr>
</tbody>
</table>
Description

By default, Dakota’s evaluation cache and restart capabilities are based on strict binary equality. This provides a performance advantage, as it permits a hash-based data structure to be used to search the evaluation cache. However, deactivating strict equality may prevent cache misses, which can occur when attempting to use a restart file on a machine different from the one on which it was generated.

7.5.5.3.1 cache_tolerance

- Keywords Area
- interface
- deactivate
- strict_cache_equality
- cache_tolerance

Specify tolerance when identifying duplicate function evaluations

Specification

Alias: none

Argument(s): REAL

Description

Described on parent page

7.5.5.4 restart_file

- Keywords Area
- interface
- deactivate
- restart_file

Deactivate writing to the restart file

Specification

Alias: none

Argument(s): none

Description

Eliminate the output of each new function evaluation to the binary restart file. This can increase speed and reduce disk storage requirements, but at the expense of a loss in the ability to recover and continue a run that terminates prematurely (e.g., due to a system crash or network problem).

Usage Tips

- This option is not recommended when function evaluations are costly or prone to failure.
• Using the `deactivate restart_file` specification will result in a zero length restart file with the default name `dakota.rst`.

7.5.6 batch

• Keywords Area

• interface

• batch

Perform evaluations in batches

Topics

This keyword is related to the topics:

• concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Default: sequential interface usage

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>size</td>
<td>Limit the number of evaluations in a batch</td>
</tr>
</tbody>
</table>

Description

When the optional `batch` keyword is used, Dakota performs evaluations in batches. In batch mode, Dakota writes the parameters for multiple (a batch of) evaluations to a single file, invokes the analysis driver once for the whole batch, and expects to find results for the entire batch in a single file after the analysis driver has exited. Batch mode may be useful when a user desires to take greater control over job management. For example, the analysis driver can perform the evaluations in the batch in any sequence or in concurrent sub-batches.

The names of the parameters file and results file are provided as command line arguments to the analysis driver, just as in a conventional, non-batch evaluation. By default, all currently available evaluations will be performed in a single batch, but the batch size can be limited using the `size` keyword.

Batch mode has a few important limitations.

• Only one batch at a time may be executed. Asynchronous execution of multiple concurrent batches is not supported.

• No `input_filter` or `output_filter` is permitted.

• Only one `analysis_driver` is allowed.

• `failure_capture` modes are restricted to `abort` and `recover`. 
Some of these restrictions may be lifted in future Dakota releases.

**File Formats**

A batch parameters file written by Dakota is simply a concatenated set of parameters files for the set of evaluations, either in aprepro or default Dakota format.

The batch results file is also a concatenated set of results files for the individual evaluations. However, because Dakota's results file format is not as rich as its parameters file format, evaluations in the batch results file must be separated by a line that begins with the '#' character.

The order of evaluations in the batch results file must match the order in the batch parameters file.

**Tagging and Work Directories**

When Dakota's work_directory feature is enabled, one directory is created per batch. If file_tag or directory_tag is used, parameters/results files and work directories are tagged with a batch Id, which is an integer that begins with 1 and increments for each new batch.

7.5.6.1 size

- **Keywords Area**
- interface
- batch
- size

Limit the number of evaluations in a batch

**Topics**

This keyword is related to the topics:

- concurrency_and_parallelism

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Default**: local: unlimited batch size, hybrid: zero batch size

**Description**

When batch execution is enabled, the default behavior is to add all available evaluations to the current batch. The **size** keyword limits the number of evaluations in a batch.

7.5.7 asynchronous

- **Keywords Area**
- interface
- asynchronous

Specify local evaluation or analysis concurrency
Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): none
Default: synchronous interface usage

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>evaluation_concurrency</td>
<td>Determine how many concurrent evaluations Dakota will schedule</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>local_evaluation_scheduling</td>
<td>Control how local asynchronous jobs are scheduled</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>analysis_concurrency</td>
<td>Limit the number of analysis drivers within an evaluation that Dakota will schedule</td>
</tr>
</tbody>
</table>

Description

The optional asynchronous keyword specifies use of asynchronous protocols (i.e., background system calls, nonblocking forks, POSIX threads) when evaluations or analyses are invoked. Evaluation and analysis concurrency can be independently controlled, as can the scheduling mode (static vs. dynamic) of the local evaluations.

Default Behavior

- when running Dakota on a single processor in asynchronous mode, the default concurrency of evaluations and analyses is all concurrency that is available. The evaluation_concurrency and analysis_concurrency specifications can be used to limit this concurrency in order to avoid machine overload or usage policy violation.

- when running Dakota on multiple processors in message passing mode, the default concurrency of evaluations and analyses on each of the servers is one (i.e., the parallelism is exclusively that of the message passing). With the evaluation_concurrency and analysis_concurrency specifications, a hybrid parallelism can be selected through combination of message passing parallelism with asynchronous parallelism on each server.

7.5.7.1 evaluation_concurrency

- Keywords Area
- interface
- asynchronous
• **evaluation_concurrency**

Determine how many concurrent evaluations Dakota will schedule

**Topics**

This keyword is related to the topics:

• **concurrency_and_parallelism**

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** local: unlimited concurrency, hybrid: no concurrency

**Description**

When asynchronous execution is enabled, the default behavior is to launch all available evaluations simultaneously. The `evaluation_concurrency` keyword can be used to limit the number of concurrent evaluations.

### 7.5.7.2 local_evaluation_scheduling

• **Keywords Area**  
• **interface**  
• **asynchronous**  
• **local_evaluation_scheduling**

Control how local asynchronous jobs are scheduled

**Topics**

This keyword is related to the topics:

• **concurrency_and_parallelism**

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** dynamic  
**Child Keywords:**
When performing asynchronous local evaluations, the local_evaluation_scheduling keyword controls how new evaluation jobs are dispatched when one completes.

The two options are:

- **dynamic**
- **static**

If the local_evaluation_scheduling is specified as dynamic (the default), each completed evaluation will be replaced by the next in the local evaluation queue.

If local_evaluation_scheduling is specified as static, each completed evaluation will be replaced by an evaluation number that is congruent modulo the evaluation_concurrency. This is helpful for relative node scheduling as described in dakota/share/dakota/examples/parallelism. For example, assuming only asynchronous local concurrency (no MPI), if the local concurrency is 6 and job 2 completes, it will be replaced with job 8.

For the case of hybrid parallelism, static local scheduling results in evaluation replacements that are modulo the total capacity, defined as the product of the evaluation concurrency and the number of evaluation servers. Both of these cases can result in idle processors if runtimes are non-uniform, so the default dynamic scheduling is preferred when relative node scheduling is not required.

7.5.7.2.1 dynamic

- **Keywords Area**
- **interface**
- **asynchronous**
- **local_evaluation_scheduling**
- **dynamic**

Dynamic local scheduling (sequential)

**Specification**

**Alias:** none
**Argument(s):** none

**Description**

If the local_evaluation_scheduling is specified as dynamic (the default), each completed evaluation will be replaced by the next in the local evaluation queue.
7.5.7.2.2  static

- Keywords Area
- interface
- asynchronous
- local_evaluation_scheduling
- static

Static local scheduling (tiled)

Specification

Alias: none
Argument(s): none

Description

If local_evaluation_scheduling is specified as static, each completed evaluation will be replaced by an evaluation number that is congruent modulo the evaluation_concurrency. This is helpful for relative node scheduling as described in dakota/share/dakota/examples/parallelism. For example, assuming only asynchronous local concurrency (no MPI), if the local concurrency is 6 and job 2 completes, it will be replaced with job 8.

For the case of hybrid parallelism, static local scheduling results in evaluation replacements that are modulo the total capacity, defined as the product of the evaluation concurrency and the number of evaluation servers. Both of these cases can result in idle processors if runtimes are non-uniform, so the default dynamic scheduling is preferred when relative node scheduling is not required.

7.5.7.3  analysis_concurrency

- Keywords Area
- interface
- asynchronous
- analysis_concurrency

Limit the number of analysis drivers within an evaluation that Dakota will schedule

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none
Argument(s): INTEGER
Default: local: unlimited concurrency, hybrid: no concurrency
When asynchronous execution is enabled and each evaluation involves multiple analysis drivers, then the default behavior is to launch all drivers simultaneously. The \texttt{analysis_concurrency} keyword can be used to limit the number of concurrently run drivers.

### 7.5.8 evaluation_servers

- Keywords Area
- interface
- evaluation_servers

Specify the number of evaluation servers when Dakota is run in parallel

#### Topics

This keyword is related to the topics:

- concurrency\_and\_parallelism

#### Specification

- **Alias**: none
- **Argument(s)**: INTEGER
- **Default**: automatic (see discussion)

#### Description

The optional \texttt{evaluation_servers} specification supports user override of the automatic parallel configuration for the number of evaluation servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the evaluation parallelism level. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

### 7.5.9 evaluation_scheduling

- Keywords Area
- interface
- evaluation_scheduling

Specify the scheduling of concurrent evaluations when Dakota is run in parallel

#### Specification

- **Alias**: none
- **Argument(s)**: none
- **Default**: automatic (see discussion)

#### Child Keywords:
7.5 interface

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Server Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>master</td>
<td>Specify a dedicated master partition for parallel evaluation scheduling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel evaluation scheduling</td>
</tr>
</tbody>
</table>

### Description

When Dakota is run in parallel, the partition type and scheduling for the evaluation servers are determined automatically. If these settings are undesirable, they may be overridden by the user using the `evaluation_scheduling` keyword.

#### 7.5.9.1 master

- **Keywords Area**
- **interface**
- **evaluation_scheduling**
- **master**

Specify a dedicated master partition for parallel evaluation scheduling

### Topics

This keyword is related to the topics:

- **concurrency_and_parallelism**

### Specification

**Alias:** none

**Argument(s):** none

### Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the evaluation servers. This reduces the number of processors available to create servers by 1.

#### 7.5.9.2 peer

- **Keywords Area**
- **interface**
- **evaluation_scheduling**
- **peer**
Specify a peer partition for parallel evaluation scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
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</thead>
<tbody>
<tr>
<td>Required (Choose One)</td>
<td></td>
<td>dynamic</td>
<td>Specify dynamic scheduling in a peer partition when Dakota is run in parallel.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>static</td>
<td>Specify static scheduling in a peer partition when Dakota is run in parallel.</td>
</tr>
</tbody>
</table>

Description

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to evaluation servers. The scheduling, static or dynamic, must also be specified.

7.5.9.2.1 dynamic

- Keywords Area
- interface
- evaluation_scheduling
- peer
- dynamic

Specify dynamic scheduling in a peer partition when Dakota is run in parallel.

Topics

This keyword is related to the topics:

- concurrency_and_parallelism
Specification

**Alias:** none

**Argument(s):** none

**Default:** dynamic (see discussion)

**Description**

In **dynamic** scheduling, evaluations are assigned to servers as earlier evaluations complete. Dynamic scheduling is advantageous when evaluations are of uneven duration.

7.5.9.2.2 **static**

- **Keywords Area**
- **interface**
- **evaluation_scheduling**
- **peer**
- **static**

Specify static scheduling in a peer partition when Dakota is run in parallel.

**Topics**

This keyword is related to the topics:

- **concurrency_and_parallelism**

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

In **static** scheduling, all available evaluations are assigned to servers in a predetermined fashion. Each completed evaluation is replaced with one congruent modulo the evaluation concurrency. For example, with 6 servers, eval number 2 will be replaced by eval number 8.

7.5.10 **processors_per_evaluation**

- **Keywords Area**
- **interface**
- **processors_per_evaluation**

Specify the number of processors per evaluation server when Dakota is run in parallel.
Keywords Area

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Default: automatic (see discussion)

Description

The optional processors_per_evaluation specification supports user override of the automatic parallel configuration for the number of processors in each evaluation server. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired server size at the evaluation parallelism level. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.

7.5.11 analysis_servers

- Keywords Area
- interface
- analysis_servers

Specify the number of analysis servers when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): INTEGER

Default: automatic (see discussion)

Description

The optional analysis_servers specification supports user override of the automatic parallel configuration for the number of analysis servers. That is, if the automatic configuration is undesirable for some reason, the user can enforce a desired number of partitions at the analysis parallelism level. Refer to ParallelLibrary and the Parallel Computing chapter of the Users Manual [5] for additional information.
7.5.12  analysis_scheduling

- Keywords Area
- interface
- analysis_scheduling

Specify the scheduling of concurrent analyses when Dakota is run in parallel

Topics

This keyword is related to the topics:

- concurrency_and_parallelism

Specification

Alias: none

Argument(s): none

Default: automatic (see discussion)

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Scheduling Mode (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>master</td>
<td>Specify a dedicated master partition for parallel analysis scheduling</td>
</tr>
<tr>
<td></td>
<td></td>
<td>peer</td>
<td>Specify a peer partition for parallel analysis scheduling</td>
</tr>
</tbody>
</table>

Description

When Dakota is run in parallel, the partition type for the analysis servers is determined automatically. If this setting is undesirable, it may be overridden by the user using the analysis_scheduling keyword.

7.5.12.1  master

- Keywords Area
- interface
- analysis_scheduling
- master

Specify a dedicated master partition for parallel analysis scheduling

Topics

This keyword is related to the topics:

- concurrency_and_parallelism
**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

This option overrides the Dakota parallel automatic configuration, forcing the use of a dedicated master partition. In a dedicated master partition, one processor (the "master") dynamically schedules work on the analysis servers. This reduces the number of processors available to create servers by 1.

7.5.12.2 peer

- Keywords Area  
- interface  
- analysis_scheduling  
- peer

Specify a peer partition for parallel analysis scheduling

**Topics**

This keyword is related to the topics:

- concurrency_and_parallelism

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

This option overrides the Dakota parallel automatic configuration, forcing the use of a peer partition. In a peer partition, all processors are available to be assigned to analysis servers. Note that unlike the case of evaluation_scheduling, it is not possible to specify static or dynamic.

7.6 responses

- Keywords Area  
- responses

Description of the model output data returned to Dakota upon evaluation of an interface.

**Topics**

This keyword is related to the topics:

- block
**Specification**

**Alias:** none  
**Argument(s):** none  

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>id_responses</td>
<td>Name the responses block; helpful when there are multiple</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>descriptors</td>
<td>Labels for the responses</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Response Type (Group 1)</td>
<td>objective_functions</td>
<td>Response type suitable for optimization</td>
</tr>
<tr>
<td></td>
<td></td>
<td>calibration_terms</td>
<td>Response type suitable for calibration or least squares</td>
</tr>
<tr>
<td></td>
<td></td>
<td>response_functions</td>
<td>Generic response type</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Gradient Type (Group 2)</td>
<td>no_gradients</td>
<td>Gradients will not be used</td>
</tr>
<tr>
<td></td>
<td></td>
<td>analytic_gradients</td>
<td>Analysis driver will return gradients</td>
</tr>
<tr>
<td></td>
<td></td>
<td>mixed_gradients</td>
<td>Gradients are needed and will be obtained from a mix of numerical and analytic sources</td>
</tr>
<tr>
<td></td>
<td></td>
<td>numerical_-gradients</td>
<td>Gradients are needed and will be approximated by finite differences</td>
</tr>
<tr>
<td>Required (Choose One)</td>
<td>Hessian Type (Group 3)</td>
<td>no_hessians</td>
<td>Hessians will not be used</td>
</tr>
<tr>
<td></td>
<td></td>
<td>numerical_-hessians</td>
<td>Hessians are needed and will be approximated by finite differences</td>
</tr>
</tbody>
</table>
Hessians are needed and will be approximated by secant updates (BFGS or SR1) from a series of gradient evaluations.

Hessians are needed and are available directly from the analysis driver.

Hessians are needed and will be obtained from a mix of numerical, analytic, and "quasi" sources.

Description

The responses specification in a Dakota input file indicates the types of data that can be returned by an interface when invoked during Dakota's execution. The specification includes three groups and two optional keywords.

The response type group indicates the type and number of responses expected by Dakota. It must be one of three types:

1. Optimization: objective and constraint functions
2. Calibration: calibration (least squares) terms and constraint functions
3. Uncertainty Quantification: generic response functions

The response type specified should be consistent with the iterative technique called for in the method specification. Certain general-purpose iterative techniques, such as parameter studies and design of experiments methods, can be used with any of these response types.

The gradient type group indicates the availability of first derivatives (gradient vectors) for the response functions.

The gradient specification also links back to the iterative method used. Gradients commonly are needed when the iterative study involves gradient-based optimization, local reliability analysis for uncertainty quantification, or local sensitivity analysis. They can optionally be used to build some types of surrogate models.

The Hessian type group specifies the availability of second derivatives (Hessian matrices) for the response functions.

Hessian availability for the response functions is similar to the gradient availability specifications, with the addition of support for "quasi-Hessians". The Hessian specification also links back to the iterative method in use; Hessians commonly would be used in gradient-based optimization by full Newton methods or in reliability analysis with second-order limit state approximations or second-order probability integrations.

Examples

Several examples follow. The first example shows an optimization data set containing an objective function and two nonlinear inequality constraints. These three functions have analytic gradient availability and no Hessian availability.
The next example shows a typical specification for a calibration data set. The six residual functions will have numerical gradients computed using the dakota finite differencing routine with central differences of 0.1% (plus/minus delta relative to current variables value = .001*value).

The last example shows a generic specification that could be used with a nondeterministic sampling iterator. The three response functions have no gradient or Hessian availability; therefore, only function values will be used by the iterator.

Parameter study and design of experiments iterators are not restricted in terms of the response data sets which may be catalogued; they may be used with any of the function specification examples shown above.

Theory

Responses specify the total data set that is available for use by the method over the course of iteration. This is distinguished from the data subset described by an active set vector (see Dakota File Data Formats in the Users Manual [Adams et al., 2010]) indicating the particular subset of the response data needed for a particular function evaluation. Thus, the responses specification is a broad description of the data to be used during a study whereas the active set vector indicates the subset currently needed.

7.6.1 id_responses

- Keywords Area
  - responses
  - id_responses

Name the responses block; helpful when there are multiple

Topics

This keyword is related to the topics:

- block_identifier

Specification

Alias: none

Argument(s): STRING

Default: use of last responses parsed
Description

The optional `id_responses` keyword accepts a string that uniquely identifies this responses block. A model can then use these responses by specifying the same string in its `responses_pointer` specification.

Default Behavior

If the `id_responses` specification is omitted, a particular responses specification will be used by a model only if that model does not include an `responses_pointer` and the responses block was the last (or only) one parsed.

Usage Tips

- It is a best practice to always use explicit responses IDs and pointers to avoid confusion.
- If only one responses block exists, then `id_responses` can be safely omitted from the responses block (and `responses_pointer` omitted from the model specification(s)), since there is no ambiguity.

Examples

For example, a model specification including

```plaintext
model
  responses_pointer = 'R1'
```

will link to a response set with

```plaintext
id_responses = 'R1'
```

7.6.2 descriptors

- Keywords Area
- responses
- descriptors

Labels for the responses

Specification

Alias: response_descriptors

Argument(s): STRINGLIST

Default: root strings plus numeric identifiers

Description

A list of strings which identify the responses. These are used in console and tabular output. Response descriptors are ordered by primary response functions (objective, calibration, or response functions), followed by inequality, then equality constraints, if present.

Default Behavior

The default descriptor strings use a response type-dependent root string plus a one-based numeric identifier:

- Objective functions: obj_fn_i
- Calibration terms: least_sq_term_i
• Nonlinear inequality constraints: nln_ineq_con_i
• Nonlinear equality constraints: nln_eq_con_i
• Response functions: response_fn_

Expected Output Dakota will label the various response functions in console and tabular output.

Usage Tips When specifying descriptors for scalar and/or field responses, include as many descriptors as top-level scalar + field responses, e.g.,

```plaintext
responses
  descriptors 'scalar1' 'scalar2' 'scalar3' 'field1' 'field2'
response_functions 5
  scalar_responses 3
  field_responses 2
  field_lengths 42 24
```

Dakota will append a numeric identifier for each field entry, for a total of 42 field1_j and 24 field2_k in this example.

Examples

```plaintext
responses
  descriptors 'cost' 'impact' 'safety'
objective_functions 2
  nonlinear_inequality_constraints 1
```

7.6.3 objective_functions

• Keywords Area
• responses
• objective_functions

Response type suitable for optimization

Specification

Alias: num_objective_functions

Argument(s): INTEGER

Child Keywords:

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<thead>
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<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>sense</td>
<td>Whether to minimize or maximize each objective function</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>primary_scale_types</td>
<td>How to scale each objective function</td>
</tr>
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</table>
### Keywords Area

<table>
<thead>
<tr>
<th>Optional</th>
<th>primary_scales</th>
<th>Characteristic values to scale each objective function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>weights</td>
<td>Specify weights for each objective function</td>
</tr>
<tr>
<td>Optional</td>
<td>nonlinear_-inequality_-constraints</td>
<td>Group to specify nonlinear inequality constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>nonlinear_equality-_constraints</td>
<td>Group to specify nonlinear equality constraints</td>
</tr>
<tr>
<td>Optional</td>
<td>scalar_objectives</td>
<td>Number of scalar objective functions</td>
</tr>
<tr>
<td>Optional</td>
<td>field_objectives</td>
<td>Number of field objective functions</td>
</tr>
</tbody>
</table>

#### Description

Specifies the number (1 or more) of objective functions $f_j$ returned to Dakota for use in the general optimization problem formulation:

\[
\begin{align*}
\text{minimize:} & \quad f(x) = \sum_j w_j f_j \\
\text{subject to:} & \quad x \in \mathbb{R}^n \\
& \quad g_L \leq g(x) \leq g_U \\
& \quad h(x) = h_t \\
& \quad a_L \leq A_x \leq a_U \\
& \quad A_e x = a_t \\
& \quad x_L \leq x \leq x_U
\end{align*}
\]

Unless `sense` is specified, Dakota will minimize the objective functions.

The keywords `nonlinear_inequality_constraints` and `nonlinear_equality_constraints` specify the number of nonlinear inequality constraints $g$, and nonlinear equality constraints $h$, respectively. When interfacing to external applications, the responses must be returned to Dakota in this order in the `results_file`:

1. objective functions
2. nonlinear_inequality_constraints
3. nonlinear_equality_constraints

An optimization problem’s linear constraints are provided to the solver at startup only and do not need to be included in the data returned on every function evaluation. Linear constraints are therefore specified in the `variables` block through the `linear_inequality_constraint_matrix` $A_i$ and `linear_equality_constraint_matrix` $A_e$.

Lower and upper bounds on the design variables $x$ are also specified in the `variables` block.

The optional keywords relate to scaling the objective functions (for better numerical results), formulating the problem as minimization or maximization, and dealing with multiple objective functions through weights $w$. If scaling is used, it is applied before multi-objective weighted sums are formed, so, e.g., when both weighting and characteristic value scaling are present the ultimate objective function would be:
\[ f = \sum_{j=1}^{n} w_j \frac{f_j}{s_j} \]

See Also

These keywords may also be of interest:

- calibration_terms
- response_functions
- method
- variables

7.6.3.1 sense

- Keywords Area
- responses
- objective_functions
- sense

Whether to minimize or maximize each objective function

Specification

Alias: none

Argument(s): STRINGLIST

Default: vector values = 'minimize'

Description

The sense keyword is used to declare whether each objective function should be minimized or maximized. The argument options are:

- "minimization" (can be abbreviated to "min")
- "maximization" (can be abbreviated to "max")

If a single string is specified it will apply to each objective function. Otherwise, the number of strings should either be equal to the number of objective functions (objective_functions). Thus, when scalar and/or field responses are specified, the number of sense strings must equal the number of scalars plus the number of fields, not the total elements in the fields.

7.6.3.2 primary_scale_types

- Keywords Area
- responses
- objective_functions
- primary_scale_types

How to scale each objective function
Specifically

**Alias:** objective_function_scale_types

**Argument(s):** STRINGLIST

**Default:** no scaling

**Description**

Each string in `primary_scale_types` indicates the scaling type for each objective function. They only have effect when the associated method specifies `scaling`.

The options are:

- `'value'` - characteristic value by which response functions will be divided. If this is chosen, then `primary_scales` must also be specified; `value` is assumed if scales are given without `scale_types`

- `'log'` - logarithmic scaling (can be used together with `primary_scales`, which can be helpfully used to negate values prior to log transformation)

(Automatic scaling is disallowed due to lack of bounds or targets for primary responses.)

If a single string is specified it will apply to all `objective_functions`. Otherwise, the length of `primary_scale_types` must have length equal to `objective_functions`. Thus, when scalar and/or field responses are specified, the number of `primary_scale_types` must equal the number of scalars plus the number of fields, not the total elements in the fields.

**Usage Tips:**

When scaling is active, it is applied to objective functions prior to any `weights` and multi-objective sum formation.

See the scaling information under specific methods, e.g., `method-**-scaling` for details on how to use this keyword.

7.6.3.3 `primary_scales`

- Keywords Area
- responses
- objective_functions
- primary_scales

Characteristic values to scale each objective function

**Specification**

**Alias:** objective_function_scales

**Argument(s):** REALLIST

**Default:** 1.0 (no scaling)

**Description**

Each real value in `primary_scales` is a nonzero characteristic value to be used in scaling each objective function. They only have effect when the associated method specifies `scaling`. 
This keyword is required for primary_scale_types of ‘value’ and optional for ‘log’. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative response functions.

Length: When specified, primary_scales must have length one of:

- One (the single value will be applied to each objective function); or

- Number of response groups (objective_functions), so the number of scalars plus the number of fields when fields are present; or

- Total number of response elements, so the number of scalar responses plus the sum of the lengths of the fields.

Usage Tips:

When primary_scales are specified, but not primary_scale_types, the scaling type is assumed to be ‘value’ for all primary response functions.

Use scale values of 1.0 to selectively avoid scaling a subset of response functions.

When scaling is active, it is applied to objective functions prior to any weights and multi-objective sum formation. See the equations in objective_functions.

See the scaling information under specific methods, e.g., method-*-scaling for details on how to use this keyword.

7.6.3.4 weights

- Keywords Area
- responses
- objective_functions
- weights

Specify weights for each objective function

Specification

Alias: multi_objective_weights

Argument(s): REALLIST

Default: equal weights

Description

For multi-objective optimization problems (where the number of objective functions is greater than 1), then a weights specification provides a simple weighted-sum approach to combining multiple objectives into a single objective:

\[ f = \sum_{i=1}^{n} w_i f_i \]

Length: The weights must have length equal to objective_functions. Thus, when scalar and/or field responses are specified, the number of weights must equal the number of scalars plus the number of fields, not the total elements in the fields.
**Default Behavior** If weights are not specified, then each response is given equal weighting:

\[ f = \sum_{i=1}^{n} \frac{f_i}{n} \]

where, in both of these cases, a "minimization" sense will retain a positive weighting for a minimizer and a "maximization" sense will apply a negative weighting.

**Usage Tips:**
Weights are applied as multipliers, scales as characteristic values / divisors.
When scaling is active, it is applied to objective functions prior to any weights and multi-objective sum formation. See the equations in **objective_functions**.

7.6.3.5 nonlinear_inequality_constraints

- **Keywords Area**
- **responses**
- **objective_functions**
- **nonlinear_inequality_constraints**

Group to specify nonlinear inequality constraints

**Topics**
This keyword is related to the topics:

- **nonlinear_constraints**

**Specification**

**Alias:** num_nonlinear_inequality_constraints

**Argument(s):** INTEGER

**Default:** 0

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>lower_bounds</td>
<td></td>
<td>Specify minimum values</td>
</tr>
<tr>
<td>Optional</td>
<td>upper_bounds</td>
<td></td>
<td>Specify maximum values</td>
</tr>
<tr>
<td>Optional</td>
<td>scale_types</td>
<td></td>
<td>How to scale each nonlinear constraint</td>
</tr>
<tr>
<td>Optional</td>
<td>scales</td>
<td></td>
<td>Characteristic values to scale each nonlinear constraint</td>
</tr>
</tbody>
</table>

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Description

Specifies the number of nonlinear inequality constraint functions returned by the interface. The `lower_bounds` and `upper_bounds` specifications provide the lower and upper bounds for 2-sided nonlinear inequalities of the form

$$g_l \leq g(x) \leq g_u$$

When constraint bounds are not specified, the problem is assumed to have one-sided inequalities bounded above by zero:

$$g(x) \leq 0.0.$$  

This provides backwards compatibility with previous Dakota versions.

In a user bounds specification, any upper bound values greater than `+bigRealBoundSize` (1.e+30, as defined in Minimizer) are treated as `+infinity` and any lower bound values less than `-bigRealBoundSize` are treated as `-infinity`. This feature is commonly used to drop one of the bounds in order to specify a 1-sided constraint (just as the default lower bounds drop out since `-DBL_MAX < -bigRealBoundSize`). The same approach is used for nonexistent linear inequality bounds and for nonexistent design variable bounds.

The `scale_types` and `scales` keywords are related to scaling of \(g(x)\). See the scaling information under specific methods, e.g., method-∗-scaling for details on how to use this keyword.

7.6.3.5.1 lower_bounds

- Keywords Area
- responses
- objective_functions
- nonlinear_inequality_constraints
- lower_bounds

Specify minimum values

Specification

**Alias:** nonlinear_inequality_lower_bounds

**Argument(s):** REALLIST

**Default:** vector values = -infinity

Description

Specify minimum values

7.6.3.5.2 upper_bounds

- Keywords Area
- responses
- objective_functions
- nonlinear_inequality_constraints
- upper_bounds

Specify maximum values
Specification

**Alias:** nonlinear_inequality_upper_bounds

**Argument(s):** REALLIST

**Default:** vector values = 0.

**Description**

Specify maximum values

7.6.3.5.3 scale_types

- Keywords Area
- responses
- objective_functions
- nonlinear_inequality_constraints
- scale_types

How to scale each nonlinear constraint

**Specification**

**Alias:** nonlinear_inequality_scale_types

**Argument(s):** STRINGLIST

**Default:** no scaling

**Description**

Each string in scale_types indicates the scaling type for each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

The options are:

- `value` - characteristic value by which nonlinear constraint values will be divided. If this is chosen, then scales must also be specified; `value` is assumed if scales are given without scale_types

- `auto` - automatic scaling based on bounds (inequalities) or targets (equalities)

- `log` - logarithmic scaling (can be used together with scales, which can be helpfully used to negate values prior to log transformation)

If a single string is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a string must be specified for each nonlinear inequality (equality) constraint.

**Usage Tips:**

See the scaling information under specific methods, e.g., method- scaling for details on how to use this keyword.
7.6.3.5.4 scales

- Keywords Area
- responses
- objective_functions
- nonlinear_inequality_constraints
- scales

Characteristic values to scale each nonlinear constraint

Specification

Alias: nonlinear_inequality_scales

Argument(s): REALLIST

Default: 1.0 (no scaling)

Description

Each real value in scales is a nonzero characteristic value to be used in scaling each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

This keyword is required for scale_types of ‘value’ and optional for ‘log’. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative response functions.

If a single scale value is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a value must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:

When scales are specified, but not scale_types, the scaling type is assumed to be ‘value’ for this set of nonlinear inequality (equality) constraints.

Use scale values of 1.0 to selectively avoid scaling a subset of response functions.

See the scaling information under specific methods, e.g., method-+-scaling for details on how to use this keyword.

7.6.3.6 nonlinear_equality_constraints

- Keywords Area
- responses
- objective_functions
- nonlinear_inequality_constraints

Group to specify nonlinear equality constraints

Topics

This keyword is related to the topics:

- nonlinear_constraints
Specification

**Alias:** num_nonlinear_equality_constraints

**Argument(s):** INTEGER

**Default:** 0

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Optional</td>
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<td>targets</td>
<td>Target values for the nonlinear equality constraint</td>
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<tr>
<td>Optional</td>
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<td>scale_types</td>
<td>How to scale each nonlinear constraint</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scales</td>
<td>Characteristic values to scale each nonlinear constraint</td>
</tr>
</tbody>
</table>

**Description**

Specifies the number of nonlinear equality constraint functions returned by the interface.

The `targets` specification provides the targets for nonlinear equalities of the form

\[ h(x) = h_t \]

and the defaults for the equality targets enforce a value of 0. for each constraint

\[ h(x) = 0.0 \]

The `scale_types` and `scales` keywords are related to scaling of \( h(x) \). See the scaling information under specific methods, e.g., method-\*scaling for details on how to use this keyword.

7.6.3.6.1 targets

- Keywords Area
- responses
- objective_functions
- nonlinear_equality_constraints
- targets

Target values for the nonlinear equality constraint

**Specification**

**Alias:** nonlinear_equality_targets

**Argument(s):** REALLIST

**Default:** vector values = 0. 

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Description

The targets specification provides the targets for nonlinear equalities of the form

\[ g(x) = g_t \]

and the defaults for the equality targets enforce a value of 0.0 for each constraint:

\[ g(x) = 0.0 \]

7.6.3.6.2 scale_types

- Keywords Area
- responses
- objective_functions
- nonlinear_equality_constraints
- scale_types

How to scale each nonlinear constraint

Specification

Alias: nonlinear_equality_scale_types

Argument(s): STRINGLIST

Default: no scaling

Description

Each string in scale_types indicates the scaling type for each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

The options are:

- ‘value’ - characteristic value by which nonlinear constraint values will be divided. If this is chosen, then scales must also be specified; ‘value’ is assumed if scales are given without scale_types

- ‘auto’ - automatic scaling based on bounds (inequalities) or targets (equalities)

- ‘log’ - logarithmic scaling (can be used together with scales, which can be helpfully used to negate values prior to log transformation)

If a single string is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a string must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:

See the scaling information under specific methods, e.g., method-+scaling for details on how to use this keyword.
7.6.3.3 scales

- Keywords Area
- responses
- objective_functions
- nonlinear_equality_constraints
- scales

Characteristic values to scale each nonlinear constraint

**Specification**

**Alias:** nonlinear_equality_scales  
**Argument(s):** REALLIST  
**Default:** 1.0 (no scaling)

**Description**

Each real value in scales is a nonzero characteristic value to be used in scaling each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

This keyword is required for scale_types of ‘value’ and optional for ‘log’. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative response functions.

If a single scale value is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a value must be specified for each nonlinear inequality (equality) constraint.

**Usage Tips:**

When scales are specified, but not scale_types, the scaling type is assumed to be ‘value’ for this set of nonlinear inequality (equality) constraints.

Use scale values of 1.0 to selectively avoid scaling a subset of response functions.

See the scaling information under specific methods, e.g., method-**scaling** for details on how to use this keyword.

7.6.3.7 scalar_objectives

- Keywords Area
- responses
- objective_functions
- scalar_objectives

Number of scalar objective functions

**Specification**

**Alias:** num_scalar_objectives  
**Argument(s):** INTEGER
Description

This keyword describes the number of scalar objective functions. It is meant to be used in conjunction with field-objectives, which describes the number of field objectives functions. The total number of objective functions, both scalar and field, is given by objective-functions. If only scalar objective functions are specified, it is not necessary to specify the number of scalar terms explicitly: one can simply say objective-functions = 5 and get 5 scalar objectives. However, if there are three scalar objectives and 2 field objectives, then objective-functions = 5 but scalar-objectives = 3 and field-objectives = 2.

Objective functions are responses that are used with optimization methods in Dakota. Currently, each term in a field objective is added to the total objective function presented to the optimizer. For example, if you have one field objective with 100 terms (e.g. a time-temperature trace with 100 time points and 100 corresponding temperature points), the 100 temperature values will be added to create the overall objective.

See Also

These keywords may also be of interest:

- field_objectives

7.6.3.8 field_objectives

- Keywords Area
- responses
- objective_functions
- field_objectives

Number of field objective functions

Specification

Alias: num_field_objectives

Argument(s): INTEGER

Child Keywords:

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<th>Dakota Keyword</th>
<th>Description</th>
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<td>lengths</td>
<td>Lengths of field responses</td>
</tr>
<tr>
<td>Optional</td>
<td>num_coordinates_per_field</td>
<td>num_coordinates_per_field</td>
<td>Number of independent coordinates for field responses</td>
</tr>
<tr>
<td>Optional</td>
<td>read_field_coordinates</td>
<td>read_field_coordinates</td>
<td>Add context to data: flag to indicate that field coordinates should be read</td>
</tr>
</tbody>
</table>

Description

This keyword describes the number of field objective functions. A field function is a set of related response values collected over a range of independent coordinate values which may or may not be specified by the user. For
example, voltage over time would be a field function, where voltage is the field_objective and time is the independent coordinate. Similarly, temperature over time and space would be a field response, where the independent coordinates would be both time and spatial coordinates such as (x,y) or (x,y,z), depending on the application. The main difference between scalar objectives and field objectives is that for field data, we plan to implement methods that take advantage of the correlation or relationship between the field values.

Note that if there is one field_objective, and it has length 100 (meaning 100 values), then the user’s simulation code must return 100 values. Also, if there are both scalar and field objectives, the user should specify the number of scalar objectives as scalar_objectives. If there are only field objectives, it still is necessary to specify both objective_functions = NN and field_objectives = NN, where NN is the number of field objectives.

Objective functions are responses that are used with optimization methods in Dakota. Currently, each term in a field objective is added to the total objective function presented to the optimizer. For example, if you have one field objective with 100 terms (e.g. a time-temperature trace with 100 time points and 100 corresponding temperature points), the 100 temperature values will be added to create the overall objective.

See Also

These keywords may also be of interest:

- scalar_objectives

7.6.3.8.1 lengths

- Keywords Area
- responses
- objective_functions
- field_objectives
- lengths

Lengths of field responses

Specification

Alias: none

Argument(s): INTEGERLIST

Description

This keyword describes the lengths of each field response. It is an integer vector of length field_responses. For example, if the field_responses = 2, an example would be lengths = 50 200, indicating that the first field response has 50 field elements but the second one has 200. The coordinate values (e.g. the independent variables) corresponding to these field responses are read in files labeled response_descriptor.coords.

See Also

These keywords may also be of interest:

- field_responses
### num_coordinates_per_field

- **Keywords Area**
- **responses**
- **objective_functions**
- **field_objectives**
- **num_coordinates_per_field**

Number of independent coordinates for field responses

**Specification**

**Alias:** none  
**Argument(s):** INTEGERLIST

**Description**

This keyword describes the number of independent coordinates for each field response. It is an integer vector of length `field_responses`. For example, if `field_responses = 2`, an example would be `num_coordinates_per_field = 2 1` means that the first field response has two sets of independent coordinates (perhaps `x`, `y` locations), but the second response only has one (for example, time where the field response is only dependent upon time). The actual coordinate values (e.g., the independent variables) corresponding to these field responses are defined in a file call `response_descriptor.coords`, where `response_descriptor` is the name of the individual field.

**See Also**

These keywords may also be of interest:

- `field_responses`

### read_field_coordinates

- **Keywords Area**
- **responses**
- **objective_functions**
- **field_objectives**
- **read_field_coordinates**

Add context to data: flag to indicate that field coordinates should be read

**Specification**

**Alias:** none  
**Argument(s):** none
Field coordinates specify independent variables (e.g. spatial or temporal coordinates) upon which the field depends. For example, the voltage level above might be a function of time, so time is the field coordinate. If the user has field coordinates to read, they need to specify `read_field_coordinates`. The field coordinates will then be read from a file named `response_descriptor.coords`, where `response_descriptor` is the user-provided descriptor for the field response. The number of columns in the coords file should be equal to the number of field coordinates.

7.6.4 calibration_terms

- **Keywords Area**
  - responses
  - calibration_terms

Response type suitable for calibration or least squares

**Specification**

*Alias*: least_squares_terms num_least_squares_terms

**Argument(s)**: INTEGER

**Child Keywords**:

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<tr>
<td>Optional</td>
<td></td>
<td>field_calibration_terms</td>
<td>Number of field calibration terms</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>primary_scales</td>
<td>Characteristic values to scale each calibration term</td>
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<td>Optional</td>
<td></td>
<td>weights</td>
<td>Specify weights for each objective function</td>
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<td>Optional</td>
<td>Calibration Data (Group 1)</td>
<td>calibration_data</td>
<td>Supply field or mixed field/scalar calibration data</td>
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<tr>
<td>Optional</td>
<td></td>
<td>calibration_data_file</td>
<td>Supply scalar calibration data only</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>simulation_-variance</td>
<td>Variance applied to simulation responses</td>
</tr>
</tbody>
</table>
Description

Responses for a calibration study are specified using `calibration_terms` and optional keywords for weighting/scaling, data, and constraints. In general when calibrating, Dakota automatically tunes parameters $\theta$ to minimize discrepancies or residuals between the model and the data:

$$R_i = \frac{y_i^{\text{Model}}(\theta) - y_i^{\text{Data}}}{\sigma_i}.$$  

Note that the problem specification affects what must be returned to Dakota in the `results_file`:

- If calibration data is not specified, then each of the calibration terms returned to Dakota through the interface is a residual $R_i$ to be driven toward zero.
- If calibration data is specified, then each of the calibration terms returned to Dakota must be a response $y_i^{\text{Model}}(\theta)$, which Dakota will difference with the data in the specified data file.

Constraints

(See general problem formulation at `objective_functions`). The keywords `nonlinear_inequality_constraints` and `nonlinear_equality_constraints` specify the number of nonlinear inequality constraints $g$, and nonlinear equality constraints $h$, respectively. When interfacing to external applications, the responses must be returned to Dakota in this order in the `results_file`:

1. calibration terms
2. nonlinear inequality constraints
3. nonlinear equality constraints

An optimization problem’s linear constraints are provided to the solver at startup only and do not need to be included in the data returned on every function evaluation. Linear constraints are therefore specified in the `variables` block through the `linear_inequality_constraint_matrix` $A_i$ and `linear_equality_constraint_matrix` $A_e$.

Lower and upper bounds on the design variables $x$ are also specified in the `variables` block.

Problem Transformations

Weighting or scaling calibration terms is often appropriate to account for measurement error or to condition the problem for easier solution. Weighting or scaling transformations are applied in the following order:

1. When present, observation error variance $\sigma_i$, or full covariance $\Sigma$, optionally specified through `experiment_variance_type`, is applied to residuals first:

$$R_i^{(1)} = \frac{y_i^{\text{Model}}(\theta) - y_i^{\text{Data}}}{\sigma_i}, \text{ or}$$

$$R_i^{(1)} = \Sigma^{-1/2} R = \Sigma^{-1/2} \left(y_i^{\text{Model}}(\theta) - y_i^{\text{Data}}\right),$$

resulting in the typical variance-weighted least squares formulation

$$\min_\theta R(\theta)^T \Sigma^{-1} R(\theta)$$
2. Any active scaling transformations are applied next, e.g., for characteristic value scaling:

\[ R_i^{(2)} = \frac{R_i^{(1)}}{s_i} \]

3. Finally the optional weights are applied in a way that preserves backward compatibility:

\[ R_i^{(3)} = \sqrt{w_i} R_i^{(2)} \]

so the ultimate least squares formulation, e.g., in a scaled and weighted case would be

\[ f = \sum_{i=1}^{n} w_i \left( \frac{y_{\text{Model}}(\theta) - y_{\text{Data}}}{s_i} \right)^2 \]

Note that specifying observation error variance and weights are mutually exclusive in a calibration problem.

Theory

Dakota calibration terms are typically used to solve problems of parameter estimation, system identification, and model calibration/inversion. Local least squares calibration problems are most efficiently solved using special-purpose least squares solvers such as Gauss-Newton or Levenberg-Marquardt; however, they may also be solved using any general-purpose optimization algorithm in Dakota. While Dakota can solve these problems with either least squares or optimization algorithms, the response data sets to be returned from the simulator are different when using \textit{objective_functions} versus \textit{calibration_terms}.

Least squares calibration involves a set of residual functions, whereas optimization involves a single objective function (sum of the squares of the residuals), i.e.,

\[ f = \sum_{i=1}^{n} R_i^2 = \sum_{i=1}^{n} \left( y_{\text{Model}}(\theta) - y_{\text{Data}} \right)^2 \]

where \( f \) is the objective function and the set of \( R_i \) are the residual functions, most commonly defined as the difference between a model response and data. Therefore, function values and derivative data in the least squares case involve the values and derivatives of the residual functions, whereas the optimization case involves values and derivatives of the sum of squares objective function. This means that in the least squares calibration case, the user must return each of \( n \) residuals separately as a separate calibration term. Switching between the two approaches sometimes requires different simulation interfaces capable of returning the different granularity of response data required, although Dakota supports automatic recasting of residuals into a sum of squares for presentation to an optimization method. Typically, the user must compute the difference between the model results and the observations when computing the residuals. However, the user has the option of specifying the observational data (e.g. from physical experiments or other sources) in a file.

See Also

These keywords may also be of interest:

- \text{objective\_functions}
- \text{response\_functions}

7.6.4.1 \textit{scalar\_calibration\_terms}

- Keywords Area
- responses
- \text{calibration\_terms}
• scalar_calibration_terms

Number of scalar calibration terms

Specification

**Alias:** none

**Argument(s):** INTEGER

**Description**

This keyword describes the number of scalar calibration terms. It is meant to be used in conjunction with field-calibration_terms, which describes the number of field calibration terms. The total number of calibration terms, both scalar and field, is given by calibration_terms. If only scalar calibration terms are specified, it is not necessary to specify the number of scalar terms explicitly: one can simply say calibration_terms = 5 and get 5 scalar terms. However, if there are three scalar terms and 2 field terms, then calibration_terms = 5 but scalar_calibration_terms = 3 and field_calibration_terms = 2.

Calibration terms are responses that are used with calibration methods in Dakota, such as least squares optimizers. Currently, each scalar term is added to the total sum-of-squares error function presented to the optimizer. However, each individual field value is added as well. For example, if you have one field calibration term with length 100 (e.g. a time - temperature trace with 100 time points and 100 temperature points), the 100 temperature values will be added to create the overall sum-of-squares error function used in calibration.

**See Also**

These keywords may also be of interest:

• field_calibration_terms

7.6.4.2 field_calibration_terms

• Keywords Area
  • responses
  • calibration_terms
  • field_calibration_terms

Number of field calibration terms

Specification

**Alias:** none

**Argument(s):** INTEGER

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
</table>

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This keyword describes the number of field calibration terms. A set of field calibration terms is a set of related response values collected over a range of independent coordinate values which may or may not be specified by the user. For example, voltage over time would be a field function, where voltage is the field objective and time is the independent coordinate. Similarly, temperature over time and space would be a field response, where the independent coordinates would be both time and spatial coordinates such as (x,y) or (x,y,z), depending on the application. The main difference between scalar calibration terms and field calibration terms is that for field data, we plan to implement methods that take advantage of the correlation or relationship between the field values. For example, with calibration, if we want to calibrate parameters that result in a good model fit to a time-temperature curve, we may have to do some interpolation between the experimental data and the simulation data. That capability requires knowledge of the independent coordinates.

Note that if there is one field_calibration_terms, and it has length 100 (meaning 100 values), then the user's simulation code must return 100 values. Also, if there are both scalar and field calibration, the user should specify the number of scalar terms as scalar_calibration_terms. If there are only field calibration terms, it still is necessary to specify both field_calibration_terms = NN and calibration_terms = NN, where NN is the number of field calibration terms.

Calibration terms are responses that are used with calibration methods in Dakota, such as least squares optimizers. Currently, each scalar term is added to the total sum-of-squares error function presented to the optimizer. However, each individual field value is added as well. For example, if you have one field calibration term with length 100 (e.g. a time-temperature trace with 100 time points and 100 temperature points), the 100 temperature values will be added to create the overall sum-of-squares error function used in calibration. We have an initial capability to interpolate the field data from the user's simulation to the experimental data. For example, if the user has thermocouple readings at 20 time points, it will be an experimental field response with 20 time points and 20 temperature values. Dakota takes the 100 simulation time-temperature values (from the example above) and interpolates those to the 20 experimental points, to create 20 residual terms (simulation minus experimental data points) that will be used in calibration.

See Also

These keywords may also be of interest:

- scalar_calibration_terms

7.6.4.2.1 lengths

- Keywords Area
- responses
- calibration_terms
- field_calibration_terms
- lengths

Lengths of field responses
7.6 responses

Specification

Alias: none  
Argument(s): INTEGERLIST

Description

This keyword describes the lengths of each field response. It is an integer vector of length field_responses. For example, if the field_responses = 2, an example would be lengths = 50 200, indicating that the first field response has 50 field elements but the second one has 200. The coordinate values (e.g. the independent variables) corresponding to these field responses are read in files labeled response_descriptor.coords.

See Also

These keywords may also be of interest:

- field_responses

7.6.4.2.2 num_coordinates_per_field

- Keywords Area
- responses
- calibration_terms
- field_calibration_terms
- num_coordinates_per_field

Number of independent coordinates for field responses

Specification

Alias: none  
Argument(s): INTEGERLIST

Description

This keyword describes the number of independent coordinates for each field response. It is an integer vector of length field_responses. For example, if the field_responses = 2, an example would be num_coordinates_per_field = 2 1 means that the first field response has two sets of independent coordinates (perhaps x, y locations), but the second response only has one (for example, time where the field response is only dependent upon time). The actual coordinate values (e.g. the independent variables) corresponding to these field responses are defined in a file call response_descriptor.coords, where response_descriptor is the name of the individual field.

See Also

These keywords may also be of interest:

- field_responses
7.6.4.2.3 read_field_coordinates

- Keywords Area
- responses
- calibration_terms
- field_calibration_terms
- read_field_coordinates

Add context to data: flag to indicate that field coordinates should be read

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Field coordinates specify independent variables (e.g. spatial or temporal coordinates) upon which the field depends. For example, the voltage level above might be a function of time, so time is the field coordinate. If the user has field coordinates to read, they need to specify `read_field_coordinates`. The field coordinates will then be read from a file named `response_descriptor.coords`, where `response_descriptor` is the user-provided descriptor for the field response. The number of columns in the coords file should be equal to the number of field coordinates.

7.6.4.3 primary_scales

- Keywords Area
- responses
- calibration_terms
- primary_scales

Characteristic values to scale each calibration term

**Specification**

**Alias:** calibration_term_scales least_squares_term_scales

**Argument(s):** REALLIST

**Default:** 1.0 (no scaling)

**Description**

Each real value in `primary_scales` is a nonzero characteristic value to be used in scaling each calibration term. They only have effect when the associated method specifies `scaling`.

**Length:** When specified, `primary_scales` must have length one of:

- One (the single value will be applied to each primary response); or
• Number of response groups (calibration_terms), so the number of scalars plus the number of fields when fields are present; or

• Total number of response elements, so the number of scalar responses plus the sum of the lengths of the fields.

Usage Tips:
Calibration only supports value-based scaling. Use scale values of 1.0 to selectively avoid scaling a subset of response functions.
Weights are applied as multipliers, scales as characteristic values / divisors.
When scaling is active, it is applied to calibration terms after any residual formation (accounting for experimental data and optionally measurement error covariance), and before any weights. See the equations in calibration_terms.
See the scaling information under specific methods, e.g., method-scaling for details on how to use this keyword.

7.6.4.4 weights

• Keywords Area
• responses
• calibration_terms
• weights

Specify weights for each objective function

Specification

Alias: calibration_weights least_squares_weights

Argument(s): REALLIST

Default: equal weights

Description

Specifies relative emphasis through weights (multipliers) w on residual elements:

\[ f = \sum_{i=1}^{n} w_i R_i^2 = \sum_{i=1}^{n} w_i (\hat{y}_{i,Model} - y_{i,Data})^2 \]

Length: The weights must have length equal to calibration_terms. Thus, when scalar and/or field responses are specified, the number of weights must equal the number of scalars plus the number of fields, not the total elements in the fields.

Default Behavior If weights are not specified, then each residual is given equal weighting.

Usage Tips:

Weights are applied as multipliers, scales as characteristic values / divisors.
When scaling is active, it is applied to calibration terms after any residual formation (accounting for experimental data and optionally measurement error covariance), and before any weights. See the equations in calibration_terms.
7.6.4.5 calibration_data

- Keywords Area
- responses
- calibration_terms
- calibration_data

Supply field or mixed field/scalar calibration data

Specification

Alias: none

Argument(s): none

Child Keywords:

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<th>Description of Group</th>
<th>Dakota Keyword</th>
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<td>num experiments</td>
<td>Add context to data: number of different experiments</td>
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<tr>
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<td>num_config_variables</td>
<td>num_config_variables</td>
<td>Add context to data: number of configuration variables.</td>
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<td>experiment_variance_type</td>
<td>experiment_variance_type</td>
<td>Add context to data: specify the type of experimental error</td>
</tr>
<tr>
<td>Optional</td>
<td>scalar_data_file</td>
<td>scalar_data_file</td>
<td>Specify a scalar data file to complement field data files (mixed case)</td>
</tr>
<tr>
<td>Optional</td>
<td>interpolate</td>
<td>interpolate</td>
<td>Flag to indicate interpolation of simulation values.</td>
</tr>
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</table>

Description

calibration_data specifies a keyword block that indicates that Dakota should read in experimental data for calibration. This block is primarily to support the reading of field calibration data. For simpler, scalar-only response cases, see calibration_data_file. The user will typically specify the number of experiments, num_experiments. If this is not specified, it is assumed there is only one experiment.

Up to four types of data may be read. They are read from a collection of files, one per response descriptor, per experiment. In this discussion, DESC refers to the response descriptor and NUM to the experiment number.

1. Values: The scalar or field-valued response function values, e.g., temperature values, voltage levels. These are read from files named DESC.NUM.dat (one per response descriptor, per experiment), e.g., volts-1.dat, volts.2.dat. Scalar files will contain a single value, while field files will each contain a column
of field response values. Without \texttt{interpolate} enabled, the lengths of these files must match those specified using the \texttt{responses-calibration_terms-field_calibration_terms-lengths} keyword.

2. \textbf{Coordinates:} Field coordinates specify independent variables (e.g., spatial or temporal coordinates) upon which the field depends. For example, the voltage might be a function of time, so time is the field coordinate. These are read from files named \texttt{DESC.NUM.coords}, each containing \texttt{num_coordinates_per_field} columns. The number of rows must be the same as in the values files described in the previous bullet.

3. \textbf{Variances:} If \texttt{experiment_variance_type} is specified, variance values are read from files \texttt{DESC.NUM.sigma}. Note that a single \texttt{experiment_variance_type} may be specified, or a unique \texttt{experiment_variance_type} per response descriptor (per scalar or field). If the \texttt{experiment_variance_type} is:
   - ‘scalar’: a single variance value will be read from the file.
   - ‘diagonal’ (field responses only): a column vector of variance values (length equal to the number of experimental data points) will be read from the file. The values are the variances of each field value for this descriptor.
   - ‘matrix’ (field responses only): a matrix of covariance values (square with size the number of experimental values) will be read from the file. The matrix is a full covariance matrix for the components of this field response. While covariance among entries in a field response may be specified, covariance among experiments is not permitted.

4. \textbf{Configuration variables:} specify the conditions corresponding to different experiments. When \texttt{responses-calibration_terms-calibration_data-num_config_variables} is specified, the configuration variable values for each experiment should be placed in a file named \texttt{experiment.NUM.config}, where the number of items in that config file are the \texttt{num_config_variables}. These variables are used as auxiliary state variables for the simulation (for example) and are not calibrated.

\textbf{Attention:} In versions of Dakota prior to 6.14, string-valued configuration variables were specified in data files with 0-based indices into the admissible values. As of Dakota 6.14, strings must be specified by value. For example a string-valued configuration variable for an experimental condition might appear in the file as \texttt{low_pressure vs. high_pressure}.

\textbf{Aggregating scalar data:} The above description is primarily relevant for field data (with files for field values, field coordinates, field variances). If the user also has scalar experimental data, it may be entered as described above, i.e., one file named \texttt{DESC.NUM.dat} per scalar response. However, an alternative is to provide the data for all scalar responses in aggregate in the simpler \texttt{scalar_data_file} format, with the number of rows of that file equal to the number of experiments. The scalar data file may be used in combination with the the separate field files described above.

\textbf{Interpolation:} One important feature of field data is the capability to interpolate between points in the field. For example, we may have simulation data at a set of responses \( y \) at time points \( t \): (\( t_1, y_{s1} \), \( t_2, y_{s2} \), etc. In this example, \( t \) is the independent coordinate for the simulation, and the simulation time and response points are denoted with subscripts \( s1, s2, s3, \ldots \). If the user has experimental data that is taken at different time points: (\( t_{e1}, y_{e1} \), \( t_{e2}, y_{e2} \), \ldots, it is necessary to interpolate the simulation data to provide estimates of the simulation response at the experimental time points to construct the residual terms (model - experiment) at the experimental time points. Dakota can perform 1-D interpolation. The user must specify the keyword \texttt{interpolate}, and also provide the field coordinates as well as field values for the experiment data.

If the \texttt{interpolate} keyword is not specified, Dakota will assume that the simulation field data and the experiment field data is taken at the same set of independent coordinate values and simply construct the difference between these field terms to create the set of residuals for the sum-of-squares calculation. When \texttt{interpolate} is specified, the simulation coordinates are assumed fixed and the same for each simulation. These simulation coordinates are provided in \texttt{DESC.coords}. However, the experiment coordinates for each experiment can be different, and are provided in the files numbered by experiment with the file names given by \texttt{DESC.NUM.coords}, as indicated above.

\textbf{Examples}

Consider this field example with three field-valued responses (volts as a function of 7 time points, amps as a function of 3 time points, and power as a function of 5 time points). The experiments each have one coordinate (time) per...
The experiments were conducted given values of two configuration variables. There are four experiments (note that there often is only one experiment).

```plaintext
responses
descriptors = 'volts' 'amps' 'power'
calibration_terms = 3

#define field terms
field_calibration_terms = 3
lengths = 7 3 5
num_coordinates_per_field = 1 1 1
read_field_coordinates

# specify the data
calibration_data
num_experiments = 4
num_config_variables = 2
experiment_variance_type 'diagonal' 'scalar' 'matrix'
```

The field data will be read from separate files:

1. **Values**: 7 values for voltage as a function of time will be read from each of volts.1.dat ... volts.-4.dat, while the 3 values of amperage come from amps.1.dat, ..., amps.4.dat and the 5 values of power will come from power.1.dat, ..., power.4.dat.

2. **Coordinates**: The corresponding time values will be read from a single column vector in each of the files volts.*.coords, amps.*.coords, and power.*.coords.

3. **Variances**: Will be read per-experiment (no covariance among experiments is permitted). The variance information is depicted graphically in `experiment_variance_type`
   - Volts specifies 'diagonal' covariance, so a column vector of length 7 with the diagonal variances will be read from each of volts.1.sigma, ..., volts.4.sigma.
   - Amps specifies 'scalar' covariance, so a single variance value will read from each of amps.1.sigma, ..., amps.4.sigma.
   - Power specifies 'matrix' covariance, so a square 5 x 5 covariance matrix will be read from each of power.1.sigma, ..., power.4.sigma.

4. **Configuration variables**: Will be read from a set of files named `experiment.1.config`, ..., `experiment.4.config`.

7.6.4.5.1 `num_experiments`

- **Keywords Area**
- **responses**
- **calibration_terms**
- **calibration_data**
- **num_experiments**

Add context to data: number of different experiments

**Specification**

**Alias**: none

**Argument(s)**: INTEGER

**Default**: 1
7.6 responses

Description

The number of different experiments. Dakota will expand the total number of residual terms based on the number of calibration terms and the number of experiments. For example, if the number of calibration terms are five scalars, and there are three experiments, the total number of residuals in the least squares formulation will be 15. See calibration_data or calibration_data_file.

7.6.4.5.2 num_config_variables

• Keywords Area
• responses
• calibration_terms
• calibration_data
• num_config_variables

Add context to data: number of configuration variables.

Specification

Alias: none
Argument(s): INTEGER
Default: 0

Description

If there are multiple experiments, there can be different configuration variables (e.g. experimental settings, boundary conditions, etc.) per experiment. See calibration_data or calibration_data_file.

During calibration, configuration variables are state variables which will be passed to the simulation, and are not treated as calibration parameters.

7.6.4.5.3 experiment_variance_type

• Keywords Area
• responses
• calibration_terms
• calibration_data
• experiment_variance_type

Add context to data: specify the type of experimental error

Specification

Alias: variance_type
Argument(s): STRINGLIST
Default: none
There are four options for specifying the experimental error (e.g. the measurement error in the data you provide for calibration purposes): 'none' (default), 'scalar', 'diagonal', or 'matrix.'

If the user specifies scalar, they can provide a scalar variance per calibration term. Note that for scalar calibration terms, only 'none' or 'scalar' are options for the measurement variance. However, for field calibration terms, there are two additional options. 'diagonal' allows the user to provide a vector of measurement variances (one for each term in the calibration field). This vector corresponds to the diagonal of the full covariance matrix of measurement errors. If the user specifies 'matrix', they can provide a full covariance matrix (not just the diagonal terms), where each element (i,j) of the covariance matrix represents the covariance of the measurement error between the i-th and j-th field values.

Usage Tips

Variance information is specified on a per-response group (descriptor), per-experiment basis. Off-diagonal covariance between response groups or between experiments is not supported.

Examples

The figure below shows an observation vector with 5 responses; 2 scalar + 3 field (each field of length \( > 1 \)). The corresponding covariance matrix has scalar variances \( \sigma_1^2, \sigma_2^2 \) for each of the scalars \( s1, s2 \), diagonal covariance \( D_3 \) for field \( f3 \), scalar covariance \( \sigma_4^2 \) for field \( f4 \), and full matrix covariance \( C_5 \) for field \( f5 \). In total, Dakota supports block diagonal covariance \( \Sigma \) across the responses, with blocks \( \Sigma_i \), which could be fully dense within a given field response group. Covariance across the highest-level responses (off-diagonal blocks) is not supported, nor is covariance between experiments.

7.6.4.5.4 scalar_data_file

- Keywords Area
- responses
- calibration_terms
- calibration_data
- scalar_data_file

Specify a scalar data file to complement field data files (mixed case)

Specification

Alias: none

Argument(s): STRING

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional/Choose One</td>
<td>Tabular Format (Group 1)</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format for experiment data</td>
</tr>
</tbody>
</table>
### Description

When calibrating both scalar and field calibration terms, to associated experimental data, the scalar data may be provided in the file named by `scalar_data_file`. This file follows the same format as: `calibration_data_file`.

#### Default Behavior

If `scalar_data_file` is omitted, all calibration data, including for scalar responses, will be read from the generic field `calibration_data` format.

7.6.4.5.4.1 custom_annotated

- **Keywords Area**
  - responses
  - calibration_terms
  - calibration_data
  - scalar_data_file
  - custom_annotated

Selects custom-annotated tabular file format for experiment data

#### Topics

This keyword is related to the topics:

- file_formats

#### Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
A custom-annotated tabular file is a whitespace-separated text file containing experiment data, including configuration variables, observations, and/or observation errors, depending on context. For experiment import, custom-annotated allows user options for whether header row and exp_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

**Default Behavior**

By default, Dakota imports tabular experiment data files in annotated format. The `custom_annotated` keyword, followed by options can be used to select other formats.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

**Examples**

Import an experimental data file containing a header row, no leading exp_id column, and experiment data in a calibration study

```plaintext
responses
...
scalar_data_file 'shock_experiment.dat'
custom_annotated header
```

Example data file with two measured quantities, three experiments:

<table>
<thead>
<tr>
<th>% velocity</th>
<th>stress</th>
</tr>
</thead>
<tbody>
<tr>
<td>18.23</td>
<td>83.21</td>
</tr>
<tr>
<td>34.14</td>
<td>93.24</td>
</tr>
<tr>
<td>22.41</td>
<td>88.92</td>
</tr>
</tbody>
</table>

**7.6.4.5.4.2 header**

- Keywords Area
- responses
- calibration_terms
- calibration_data
- scalar_data_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file
7.6 responses

Specification

Alias: none
Argument(s): none
Default: no header

Description

See description of parent custom.annotated

7.6.4.5.4.3 exp_id

- Keywords Area
- responses
- calibration_terms
- calibration_data
- scalar_data_file
- custom.annotated
- exp_id

Enable experiment ID column in custom-annotated tabular file

Specification

Alias: none
Argument(s): none
Default: no exp_id column

Description

See description of parent custom.annotated

7.6.4.5.4.4 annotated

- Keywords Area
- responses
- calibration_terms
- calibration_data
- scalar_data_file
- annotated

Selects annotated tabular file format for experiment data
Topics

This keyword is related to the topics:

- file_formats

Specification

Alias: none
Argument(s): none
Default: annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. For experiment data files, each subsequent row contains an experiment ID, followed by data for configuration variables, observations, and/or observation errors, depending on context.

Default Behavior

By default, Dakota imports tabular experiment data files in annotated format. The annotated keyword can be used to explicitly specify this.

Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

Examples

Import an annotated experimental data file containing a header row, leading exp_id column, and experiment data in a calibration study

    responses
    ...
    scalar_data_file 'shock_experiment.dat'
    annotated

Example data file with two measured quantities, three experiments:

    %exp_id    velocity    stress
    1 18.23 83.21
    2 34.14 93.24
    3 22.41 88.92

7.6.4.5.4.5 freeform

- Keywords Area
- responses
- calibration_terms
- calibration_data
Selects free-form tabular file format for experiment data

Topics
This keyword is related to the topics:

- file_formats

Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. For experiment data files, each row contains data for configuration variables, observations, and/or observation errors, depending on context.

**Default Behavior**

By default, Dakota imports tabular experiment data files in annotated format. Specify `freeform` to instead select this format.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

**Examples**

Import a free-form experimental data file containing raw experiment data in a calibration study

```plaintext
responses
...
scalar_data_file 'shock_experiment.dat'
    freeform
```

Example data file with two measured quantities, three experiments:

```
18.23  83.21
34.14  93.24
22.41  88.92
```
7.6.4.5.5 interpolate

• Keywords Area
• responses
• calibration_terms
• calibration_data
• interpolate

Flag to indicate interpolation of simulation values.

Specification

Alias: none
Argument(s): none

Description

If `interpolate` is specified, Dakota will interpolate between the simulation data and the experiment data to calculate the residuals for calibration methods. Specifically, the simulation data is interpolated onto the experimental data points. So, if the simulation data is a field of length 100 with one independent coordinate, and the experiment data is of length 5 with one independent coordinate, the interpolation is done between the 100 \((t, f)\) simulation points (where \(t\) is the independent coordinate and \(f\) is the simulation field value) onto the five \((t_e, f_e)\) points to obtain the residual differences between the simulation and experiment. See `calibration_data`.

7.6.4.6 calibration_data_file

• Keywords Area
• responses
• calibration_terms
• calibration_data_file

Supply scalar calibration data only

Specification

Alias: least_squares_data_file
Argument(s): STRING
Default: none
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Tabular Format (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>custom_annotated</td>
<td>custom_annotated</td>
<td>Selects custom-annotated tabular file format for experiment data</td>
</tr>
<tr>
<td></td>
<td>annotated</td>
<td>annotated</td>
<td>Selects annotated tabular file format for experiment data</td>
</tr>
<tr>
<td></td>
<td>freeform</td>
<td>freeform</td>
<td>Selects free-form tabular file format for experiment data</td>
</tr>
<tr>
<td>Optional</td>
<td>num_experiments</td>
<td>Add context to data: number of different experiments</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>num_config_variables</td>
<td>Add context to data: number of configuration variables.</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td>experiment_variance_type</td>
<td>Add context to data: specify the type of experimental error</td>
<td></td>
</tr>
</tbody>
</table>

Description

Enables text file import of experimental observations for use in calibration, for scalar responses only, with optional scalar variance information. For more complex data import cases see calibration_data Dakota will calibrate model variables to best match these data.

Key options include:

- **format**: whether the data file is in annotated, custom_annotated, or freeform format
- **content**: where num_experiments, num_config_variables, and experiment_variance_type indicate which columns appear in the data.

In the most general case, the content of the data file is described by the arguments of three optional parameters.

- **num_experiments** ($N_{exp}$)
  Default: $N_{exp} = 1$
  This indicates that the data represent multiple experiments, where each experiment might be conducted with different values of configuration variables. An experiment can also be thought of as a replicate, where the experiments are run at the same values of the configuration variables.

- **num_config_variables** ($N_{cfg}$)
  Configuration variables specify the values of experimental conditions at which data were collected. The variables in these columns must correspond to state variables in the calibration study. The simulation model will be run at each configuration and compared to the appropriate experiment data.

- **experiment_variance_type** (‘none’ or ‘scalar’)
  This indicates if the data file contains variances for measurement error of the experimental data. The default is ‘none’.
While some components may be omitted, the most complete version of an annotated calibration data file could include columns corresponding to experiment ID, configuration variables, function value observations, and variances (observation errors), shown here in annotated format:

```
exp_id | configuration xvars | y data observations | y data variances
1 7.8 7 | 21.9372 1.8687 | 0.25 0.04
2 8.6 2 | 19.0779 4.8976 | 0.25 0.04
3 8.4 8 | 38.2758 4.4559 | 0.25 0.04
4 4.2 1 | 39.7600 6.4631 | 0.25 0.04
```

Each row in the file corresponds to an experiment or replicate observation of an experiment to be compared to the model output. This example shows 4 experiments, governed by two configuration variables (one real-valued and one integer-valued), two responses (QOIs), and corresponding observation errors with standard deviation 0.5 and 0.2.

**Usage Tips**

- The `calibration_data_file` keyword is used when only scalar calibration terms are present. If there are field calibration terms, instead use `calibration_data`. For mixed scalar and field calibration terms, one may use the `scalar_data_file` specification, which uses the format described on this page.

- **Attention:** In versions of Dakota prior to 6.14, string-valued configuration variables were specified in data files with 0-based indices into the admissible values. As of Dakota 6.14, strings must be specified by value. For example a string-valued configuration variable for an experimental condition might appear in the file as `low_pressure` vs. `high_pressure`.

**Examples**

**Simple Case:** In the simplest case, no data content descriptors are specified:

```
responses
calibration_terms = 2
descriptors = 'volts' 'amps'
calibration_data_file = 'circuit.dat'
annotated
```

And the data file `circuit.dat` must contain only the `y Data` observations which represent a single experimental observation. In this case, the data file should have `Nterms = 2` columns (for volts, amps) and 1 row, where `Nterms` is the value of `calibration_terms`. The data file is shown here in annotated format:

```
exp_id | y data observations
1 21.9372 1.8687
```

For each function evaluation, Dakota will run the analysis driver, which must return `Nterms = 2` model responses. Then the residuals are computed as:

```
R_i = y_i^{Model} - y_i^{Data}.
```

These residuals can be weighted using `weights`.

**Multiple experiments:** One might specify `num_experiments N_e` indicating that there are multiple experiments. When multiple experiments are present, Dakota will expand the number of residuals for the repeat measurement data and difference with the data accordingly. For example, if the user has `N_e = 4` experiments in the example above with 2 calibration terms, the input file would contain

```
responses
calibration_terms = 2
descriptors = 'volts' 'amps'
calibration_data_file = 'circuit.dat'
annotated
calibration_data_file = 'circuit.dat'
```

Dakota Version 6.15 Reference Manual generated on November 5, 2021
And the `calibration_data_file` would need to contain 2 rows (one for each experiment), and each row should contain 2 experimental data values that will be differenced with respect to the appropriate model response:

```
%exp_id | y data observations
1      | 21.9372 1.8687
2      | 19.0779 4.8976
3      | 38.2758 4.4559
4      | 39.7600 6.4631
```

To summarize, Dakota will calculate the sum of the squared residuals as:

\[ f = \sum_{i=1}^{N_{\text{terms}}} R_i^2 \]

where the residuals now are calculated as:

\[ R_i = y_{\text{Model}}(\theta) - y_{\text{Data}}. \]

**With experimental variances:** If information is known about the measurement error and the uncertainty in the measurement, that can be specified by sending the measurement error variance to Dakota. In this case, the keyword `experiment_variance_type` is added, followed by a string of variance types of length one or of length `N_{\text{terms}}`, where `N_{\text{terms}}` is the value of `calibration_terms`. The `experiment_variance_type` for each response can be ‘none’ or ‘scalar’. NOTE: you must specify the same `experiment_variance_type` for all scalar terms. That is, they will all be ‘none’ or all be ‘scalar.’

```python
responses
  calibration_terms = 2
  descriptors = ‘volts’ ‘amps’
  calibration_data_file = ‘circuit.dat’
  annotated
  experiment_variance_type ‘scalar’
```

For each response that has a ‘scalar’ variance type, each row of the datafile will now have `N_{\text{terms}} = 2` of `y` data values (volts, amps) followed by `N_{\text{terms}} = 2` columns that specify the measurement error (in units of variance, not standard deviation) for volts, amps. An example with two experiments in annotated format:

```
%exp_id | y data observations | y data variances
1      | 21.9372 1.8687      | 0.25 0.04
2      | 19.0779 4.8976      | 0.25 0.04
3      | 38.2758 4.4559      | 0.25 0.04
4      | 39.7600 6.4631      | 0.25 0.04
```

Dakota will run the analysis driver, which must return `N_{\text{terms}}` responses. Then the residuals are computed as:

\[ R_i = \frac{y_{\text{Model}} - y_{\text{Data}}}{\sqrt{\text{var}_i}} \]

for \( i = 1 \ldots N_{\text{terms}} \).

**Putting all the options together:** Specifying all these options together might look like

```python
responses
  calibration_terms = 2
  descriptors = ‘volts’ ‘amps’
  calibration_data_file = ‘circuit.dat’
  annotated
  num_experiments = 4
  experiment_variance_type ‘scalar’
```

Dakota will expect a data file

```
%exp_id | configuration xvars | y data observations | y data variances
1      | 7.8 7               | 21.9372 1.8687     | 0.25 0.04
2      | 8.6 2               | 19.0779 4.8976     | 0.25 0.04
3      | 8.4 8               | 38.2758 4.4559     | 0.25 0.04
4      | 4.2 1               | 39.7600 6.4631     | 0.25 0.04
```
To compute residuals for each experiment, e.g., exp_id = 4, Dakota will

1. Evaluate the computational model at the specified configuration (state variables = [4.2, 1]).
2. Difference the resulting 2 function values with the data [39.7600 volts, 6.4631 amps]
3. Weight by the standard deviation = sqrt([0.25 0.04])

See Also

These keywords may also be of interest:

• calibration_data

7.6.4.6.1 custom_annotated

• Keywords Area
• responses
• calibration_terms
• calibration_data_file
• custom_annotated

Selects custom-annotated tabular file format for experiment data

Topics

This keyword is related to the topics:

• file_formats

Specification

Alias: none

Argument(s): none

Default: annotated format

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>header</td>
<td></td>
<td>Enable header row in custom-annotated tabular file</td>
</tr>
</tbody>
</table>
### Description

A custom-annotated tabular file is a whitespace-separated text file containing experiment data, including configuration variables, observations, and/or observation errors, depending on context. For experiment import, custom-annotated allows user options for whether header row and exp_id column appear in the tabular file, thus bridging freeform and (fully) annotated.

### Default Behavior

By default, Dakota imports tabular experiment data files in annotated format. The `custom_annotated` keyword, followed by options can be used to select other formats.

### Usage Tips

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.

- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

### Examples

**Import an experimental data file containing a header row, no leading exp_id column, and experiment data in a calibration study**

```plaintext
responses
  ...
scalar_data_file 'shock_experiment.dat'
custom_annotated header
```

**Example data file with two measured quantities, three experiments:**

```plaintext
% velocity  stress
18.23      83.21
34.14      93.24
22.41      88.92
```

### 7.6.4.6.1.1 header

- Keywords Area
- responses
- calibration_terms
- calibration_data_file
- custom_annotated
- header

Enable header row in custom-annotated tabular file
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no header

Description

**See description of parent** custom.annotated

7.6.4.6.1.2  exp_id

- Keywords Area
- responses
- calibration_terms
- calibration_data_file
- custom.annotated
- exp_id

Enable experiment ID column in custom-annotated tabular file

Specification

**Alias:** none  
**Argument(s):** none  
**Default:** no exp_id column

Description

**See description of parent** custom.annotated

7.6.4.6.2  annotated

- Keywords Area
- responses
- calibration_terms
- calibration_data_file
- annotated

Selects annotated tabular file format for experiment data

Topics

This keyword is related to the topics:

- file_formats
Specification

**Alias:** none

**Argument(s):** none

**Default:** annotated format

Description

An annotated tabular file is a whitespace-separated text file with one leading header row of comments/column labels. For experiment data files, each subsequent row contains an experiment ID, followed by data for configuration variables, observations, and/or observation errors, depending on context.

**Default Behavior**

By default, Dakota imports tabular experiment data files in annotated format. The `annotated` keyword can be used to explicitly specify this.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

**Examples**

Import an annotated experimental data file containing a header row, leading exp_id column, and experiment data in a calibration study

```dakota
responses
... scalar_data_file 'shock_experiment.dat'
annotated
```

Example data file with two measured quantities, three experiments:

```
%exp_id    velocity    stress
1          18.23       83.21
2          34.14       93.24
3          22.41       88.92
```

**7.6.4.6.3 freeform**

- **Keywords Area**
- **responses**
- **calibration_terms**
- **calibration_data_file**
- **freeform**

Selects free-form tabular file format for experiment data

**Topics**

This keyword is related to the topics:

- **fileFormats**
Specification

**Alias:** none  
**Argument(s):** none  
**Default:** annotated format

**Description**

A freeform tabular file is whitespace-separated text file with no leading header row and no leading columns. For experiment data files, each row contains data for configuration variables, observations, and/or observation errors, depending on context.

**Default Behavior**

By default, Dakota imports tabular experiment data files in annotated format. Specify `freeform` to instead select this format.

**Usage Tips**

- Prior to October 2011, calibration and surrogate data files were in free-form format. They now default to annotated format, though freeform remains an option.
- When importing tabular data, a warning will be generated if a specific number of data are expected, but extra is found and an error generated when there is insufficient data.

**Examples**

Import a free-form experimental data file containing raw experiment data in a calibration study

```plaintext
responses ...
scalar_data_file 'shock_experiment.dat'
freeform
```

Example data file with two measured quantities, three experiments:

```
18.23   83.21  
34.14   93.24  
22.41   88.92  
```

7.6.4.6.4 num_experiments

- **Keywords Area**
- **responses**
- **calibration_terms**
- **calibration_data_file**
- **num_experiments**

Add context to data: number of different experiments

**Specification**

**Alias:** none  
**Argument(s):** INTEGER  
**Default:** 1
Description

The number of different experiments. Dakota will expand the total number of residual terms based on the number of calibration terms and the number of experiments. For example, if the number of calibration terms are five scalars, and there are three experiments, the total number of residuals in the least squares formulation will be 15. See calibration_data or calibration_data_file.

7.6.4.6.5 num_config_variables

- Keywords Area
- responses
- calibration_terms
- calibration_data_file
- num_config_variables

Add context to data: number of configuration variables.

Specification

Alias: none

Argument(s): INTEGER

Default: 0

Description

If there are multiple experiments, there can be different configuration variables (e.g. experimental settings, boundary conditions, etc.) per experiment. See calibration_data or calibration_data_file.

During calibration, configuration variables are state variables which will be passed to the simulation, and are not treated as calibration parameters.

7.6.4.6.6 experiment_variance_type

- Keywords Area
- responses
- calibration_terms
- calibration_data_file
- experiment_variance_type

Add context to data: specify the type of experimental error

Specification

Alias: variance_type

Argument(s): STRINGLIST

Default: none
Description

There are four options for specifying the experimental error (e.g. the measurement error in the data you provide for calibration purposes): 'none' (default), 'scalar', 'diagonal', or 'matrix.'

If the user specifies scalar, they can provide a scalar variance per calibration term. Note that for scalar calibration terms, only 'none' or 'scalar' are options for the measurement variance. However, for field calibration terms, there are two additional options. ‘diagonal’ allows the user to provide a vector of measurement variances (one for each term in the calibration field). This vector corresponds to the diagonal of the full covariance matrix of measurement errors. If the user specifies 'matrix', they can provide a full covariance matrix (not just the diagonal terms), where each element (i,j) of the covariance matrix represents the covariance of the measurement error between the i-th and j-th field values.

Usage Tips

Variance information is specified on a per-response group (descriptor), per-experiment basis. Off-diagonal covariance between response groups or between experiments is not supported.

Examples

The figure below shows an observation vector with 5 responses; 2 scalar + 3 field (each field of length > 1). The corresponding covariance matrix has scalar variances $\sigma_1^2, \sigma_2^2$ for each of the scalars $s1, s2$, diagonal covariance $D_3$ for field $f3$, scalar covariance $\sigma_4^2$ for field $f4$, and full matrix covariance $C_5$ for field $f5$. In total, Dakota supports block diagonal covariance $\Sigma$ across the responses, with blocks $\Sigma_i$, which could be fully dense within a given field response group. Covariance across the highest-level responses (off-diagonal blocks) is not supported, nor is covariance between experiments.

7.6.4.7 simulation_variance

- **Keywords Area**
- responses
- calibration_terms
- simulation_variance

Variance applied to simulation responses

Specification

**Alias:** none

**Argument(s):** REALLIST

**Default:** no variance

Description

The variance that is applied to simulations run by Dakota, i.e. using the `analysis_drivers` command. The user may supply a single variance or a vector of variances of length equal to the number of responses. In both cases, the values provided are treated as scalar variance types. If a single variance is provided, it is applied to all responses produced by the simulation code. If a vector is provided, each variance is applied to the corresponding response output produced by the simulation code.

It is important to note that the variance defined by this keyword differs from that defined using `experiment_variance_type`. These two commands apply to user-provided calibration data, specified, for example, by `calibration_data` or `calibration_data_file`. However, `simulation_variance` applies to those responses produced by simulation code that is run by Dakota, as described above.
Usage Tips
Currently, this keyword is only in use as part of the algorithm implemented by experimental_design. In this algorithm, two models (usually, one high-fidelity and one low-fidelity) are provided to Dakota, each with their own responses section of the input script, and each of which is allowed its own simulation_variance. The variance specified in the responses block belonging to the high-fidelity model is applied to any new high-fidelity data that is produced by Dakota running the high-fidelity model. In the experimental_design algorithm, low-fidelity model responses are used during the calibration of the model parameters, the calculation of the mutual information, and the calculation of any posterior statistics after the algorithm is complete. The simulation_variance is applied to the low-fidelity model responses that are used in the calculation of the mutual information. See the User's and Theory Manuals for more information.

Examples
The example below shows two responses blocks, one for the low-fidelity model and one for the high-fidelity model. Both contain simulation_variance commands that will apply to the low- and high-fidelity model responses, respectively.

```
responses,
    id_responses = 'low-fidelity'
    calibration_terms = 1
    simulation_variance = 0.5

responses,
    id_responses = 'high-fidelity'
    calibration_terms = 1
    calibration_data_file = 'dakota_bayes_expdesign.dat'
    freeform
    num_config_variables = 1
    num_experiments = 1
    experiment_variance_type = 'none'
    simulation_variance = 1.2
```

7.6.4.8 nonlinear_inequality_constraints

- Keywords Area
- responses
- calibration_terms
- nonlinear_inequality_constraints

Group to specify nonlinear inequality constraints

Topics
This keyword is related to the topics:

- nonlinear_constraints

Specification
Alias: num_nonlinear_inequality_constraints
Argument(s): INTEGER
Default: 0
Child Keywords:
<table>
<thead>
<tr>
<th>Required/Optional Group</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>lower_bounds</td>
<td>Specify minimum values</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>upper_bounds</td>
<td>Specify maximum values</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scale_types</td>
<td>How to scale each nonlinear constraint</td>
<td></td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>scales</td>
<td>Characteristic values to scale each nonlinear constraint</td>
<td></td>
</tr>
</tbody>
</table>

**Description**

Specifies the number of nonlinear inequality constraint functions returned by the interface.

The `lower_bounds` and `upper_bounds` specifications provide the lower and upper bounds for 2-sided nonlinear inequalities of the form

\[ g_l \leq g(x) \leq g_u \]

When constraint bounds are not specified, the problem is assumed to have one-sided inequalities bounded above by zero:

\[ g(x) \leq 0.0. \]

This provides backwards compatibility with previous Dakota versions.

In a user bounds specification, any upper bound values greater than `+bigRealBoundSize` (1.e+30, as defined in Minimizer) are treated as `+infinity` and any lower bound values less than `-bigRealBoundSize` are treated as `-infinity`. This feature is commonly used to drop one of the bounds in order to specify a 1-sided constraint (just as the default lower bounds drop out since `-DBL_MAX < -bigRealBoundSize`). The same approach is used for nonexistent linear inequality bounds and for nonexistent design variable bounds.

The `scale_types` and `scales` keywords are related to scaling of \( g(x) \). See the scaling information under specific methods, e.g., method-`+`-scaling for details on how to use this keyword.

7.6.4.8.1 lower_bounds

- **Keywords Area**
- **responses**
- **calibration_terms**
- **nonlinear_inequality_constraints**
- **lower_bounds**

Specify minimum values

**Specification**

**Alias:** nonlinear_inequality_lower_bounds

**Argument(s):** REALLIST

**Default:** vector values = -infinity
Description

Specify minimum values

7.6.4.8.2  upper_bounds

- Keywords Area
- responses
- calibration_terms
- nonlinear_inequality_constraints
- upper_bounds

Specify maximum values

Specification

Alias: nonlinear_inequality_upper_bounds
Argument(s): REALLIST
Default: vector values = 0.

Description

Specify maximum values

7.6.4.8.3  scale_types

- Keywords Area
- responses
- calibration_terms
- nonlinear_inequality_constraints
- scale_types

How to scale each nonlinear constraint

Specification

Alias: nonlinear_inequality_scale_types
Argument(s): STRINGLIST
Default: no scaling

Description

Each string in scale_types indicates the scaling type for each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

The options are:
'value' - characteristic value by which nonlinear constraint values will be divided. If this is chosen, then scales must also be specified; 'value' is assumed if scales are given without scale_types.

'auto' - automatic scaling based on bounds (inequalities) or targets (equalities).

'log' - logarithmic scaling (can be used together with scales, which can be helpfully used to negate values prior to log transformation).

If a single string is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a string must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:
See the scaling information under specific methods, e.g., method- scaling for details on how to use this keyword.

7.6.4.8.4 scales

Characteristics values to scale each nonlinear constraint

Specification

Alias: nonlinear_inequality_scales

Argument(s): REALLIST

Default: 1.0 (no scaling)

Description

Each real value in scales is a nonzero characteristic value to be used in scaling each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

This keyword is required for scale_types of 'value' and optional for 'log'. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative response functions.

If a single scale value is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a value must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:

When scales are specified, but not scale_types, the scaling type is assumed to be 'value' for this set of nonlinear inequality (equality) constraints.

Use scale values of 1.0 to selectively avoid scaling a subset of response functions.

See the scaling information under specific methods, e.g., method- scaling for details on how to use this keyword.
Group to specify nonlinear equality constraints

Topics

This keyword is related to the topics:

- nonlinear_constraints

Specification

Alias: num_nonlinear_equality_constraints

Argument(s): INTEGER

Default: 0

Child Keywords:

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td>targets</td>
<td></td>
<td>Target values for the nonlinear equality constraint</td>
</tr>
<tr>
<td>Optional</td>
<td>scale_types</td>
<td></td>
<td>How to scale each nonlinear constraint</td>
</tr>
<tr>
<td>Optional</td>
<td>scales</td>
<td></td>
<td>Characteristic values to scale each nonlinear constraint</td>
</tr>
</tbody>
</table>

Description

Specifies the number of nonlinear equality constraint functions returned by the interface. The targets specification provides the targets for nonlinear equalities of the form

\[ h(x) = h_i \]

and the defaults for the equality targets enforce a value of 0. for each constraint

\[ h(x) = 0.0 \]

The scale_types and scales keywords are related to scaling of \( h(x) \). See the scaling information under specific methods, e.g., method-++-scaling for details on how to use this keyword.
• targets

Target values for the nonlinear equality constraint

Specification

Alias: nonlinear_equality_targets
Argument(s): REALLIST
Default: vector values = 0.

Description

The targets specification provides the targets for nonlinear equalities of the form
\[ g(x) = g_t \]
and the defaults for the equality targets enforce a value of 0.0 for each constraint:
\[ g(x) = 0.0 \]

7.6.4.9.2 scale_types

• Keywords Area
• responses
• calibration_terms
• nonlinear_equality_constraints
• scale_types

How to scale each nonlinear constraint

Specification

Alias: nonlinear_equality_scale_types
Argument(s): STRINGLIST
Default: no scaling

Description

Each string in scale_types indicates the scaling type for each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

The options are:

• 'value' - characteristic value by which nonlinear constraint values will be divided. If this is chosen, then scales must also be specified; 'value' is assumed if scales are given without scale_types

• 'auto' - automatic scaling based on bounds (inequalities) or targets (equalities)
• ‘log’ - logarithmic scaling (can be used together with scales, which can be helpfully used to negate values prior to log transformation)

If a single string is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a string must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:
See the scaling information under specific methods, e.g., method-<scaling for details on how to use this keyword.

7.6.4.9.3 scales

- Keywords Area
- responses
- calibration_terms
- nonlinear_equality_constraints
- scales

Characteristic values to scale each nonlinear constraint

Specification

Alias: nonlinear_equality_scales
Argument(s): REALLIST
Default: 1.0 (no scaling)

Description

Each real value in scales is a nonzero characteristic value to be used in scaling each nonlinear inequality (equality) constraint. They only have effect when the associated method specifies scaling.

This keyword is required for scale_types of ‘value’ and optional for ‘log’. When specified in conjunction with log, scale values are applied prior to the logarithm, to permit log scaling of strictly negative response functions.

If a single scale value is specified it will apply to all of the nonlinear inequality (equality) constraints. Otherwise a value must be specified for each nonlinear inequality (equality) constraint.

Usage Tips:

When scales are specified, but not scale_types, the scaling type is assumed to be ‘value’ for this set of nonlinear inequality (equality) constraints.

Use scale values of 1.0 to selectively avoid scaling a subset of response functions.

See the scaling information under specific methods, e.g., method-<scaling for details on how to use this keyword.

7.6.5 response_functions

- Keywords Area
- responses
- response_functions

Generic response type
Specification

**Alias**: num_response_functions  
**Argument(s)**: INTEGER

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>scalar_responses</td>
<td>Number of scalar response functions</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>field_responses</td>
<td>Number of field responses functions</td>
</tr>
</tbody>
</table>

Description

A generic response data set is specified using `response_functions`. Each of these functions is simply a response quantity of interest with no special interpretation taken by the method in use.

Whereas objective, constraint, and residual functions have special meanings for optimization and least squares algorithms, the generic response function data set need not have a specific interpretation and the user is free to define whatever functional form is convenient.

Theory

This type of data set is used by uncertainty quantification methods, in which the effect of parameter uncertainty on response functions is quantified, and can also be used in parameter study and design of experiments methods (although these methods are not restricted to this data set), in which the effect of parameter variations on response functions is evaluated.

See Also

These keywords may also be of interest:

- `objective_functions`
- `calibration_terms`

7.6.5.1 scalar_responses

- Keywords Area
- responses
- response_functions
- scalar_responses

Number of scalar response functions

Specification

**Alias**: num_scalar_responses  
**Argument(s)**: INTEGER
Description

This keyword describes the number of scalar response functions. It is meant to be used in conjunction with `field_responses`, which describes the number of field response functions. The total number of response functions, both scalar and field, is given by `response_functions`. If only scalar responses functions are specified, it is not necessary to specify the number of scalar terms explicitly: one can simply say `response_functions = 5` and get 5 scalar responses. However, if there are three scalar responses and 2 field responses, then `response_functions = 5` but `scalar_responses = 3` and `field_responses = 2`.

This type of data set is used by uncertainty quantification methods, in which the effect of parameter uncertainty on response functions is quantified, and can also be used in parameter study and design of experiments methods (although these methods are not restricted to this data set), in which the effect of parameter variations on response functions is evaluated.

See Also

These keywords may also be of interest:

- `field_responses`

7.6.5.2 field_responses

- Keywords Area
- responses
- response_functions
- field_responses

Number of field responses functions

Specification

Alias: `num_field_responses`

Argument(s): INTEGER

Child Keywords:

<table>
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<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tr>
<td>Required</td>
<td>lengths</td>
<td></td>
<td>Lengths of field responses</td>
</tr>
<tr>
<td>Optional</td>
<td>num_coordinates_per_field</td>
<td></td>
<td>Number of independent coordinates for field responses</td>
</tr>
<tr>
<td>Optional</td>
<td>read_field_coordinates</td>
<td></td>
<td>Add context to data: flag to indicate that field coordinates should be read</td>
</tr>
</tbody>
</table>

Description

This keyword describes the number of field response functions. A field function is a set of related response values collected over a range of independent coordinate values which may or may not be specified by the user. For
example, voltage over time would be a field function, where voltage is the field_objective and time is the independent coordinate. Similarly, temperature over time and space would be a field response, where the independent coordinates would be both time and spatial coordinates such as (x,y) or (x,y,z), depending on the application. The main difference between scalar responses and field responses is that for field data, we plan to implement methods that take advantage of the correlation or relationship between the field values.

Note that if there is one field_response, and it has length 100 (meaning 100 values), then the user’s simulation code must return 100 values. Also, if there are both scalar and field responses, the user should specify the number of scalar responses as scalar_responses. If there are only field responses, it is still necessary to specify both response_functions = NN and field_responses = NN, where NN is the number of field responses.

This type of data set is used by uncertainty quantification methods, in which the effect of parameter uncertainty on response functions is quantified, and can also be used in parameter study and design of experiments methods (although these methods are not restricted to this data set), in which the effect of parameter variations on response functions is evaluated. Currently, field response functions will be translated back to scalar responses. So, a field of length 100 will be treated as 100 separate scalar responses. However, in future versions of Dakota, we plan to implement methods which can exploit the nature of field data.

See Also

These keywords may also be of interest:

- scalar_responses

7.6.5.2.1 lengths

- Keywords Area
- responses
- response_functions
- field_responses
- lengths

Lengths of field responses

Specification

Alias: none

Argument(s): INTEGERLIST

Description

This keyword describes the lengths of each field response. It is an integer vector of length field_responses. For example, if the field_responses = 2, an example would be lengths = 50 200, indicating that the first field response has 50 field elements but the second one has 200. The coordinate values (e.g. the independent variables) corresponding to these field responses are read in files labeled response_descriptor.coords.

See Also

These keywords may also be of interest:

- field_responses
7.6.5.2.2 num_coordinates_per_field

- Keywords Area
- responses
- response_functions
- field_responses
- num_coordinates_per_field

Number of independent coordinates for field responses

**Specification**

**Alias:** none

**Argument(s):** INTEGERLIST

**Description**

This keyword describes the number of independent coordinates for each field response. It is an integer vector of length field_responses. For example, if the field_responses = 2, an example would be num_coordinates_per_field = 2 1 means that the first field response has two sets of independent coordinates (perhaps x, y locations), but the second response only has one (for example, time where the field response is only dependent upon time). The actual coordinate values (e.g. the independent variables) corresponding to these field responses are defined in a file call response_descriptor.coords, where response_descriptor is the name of the individual field.

**See Also**

These keywords may also be of interest:

- field_responses

7.6.5.2.3 read_field_coordinates

- Keywords Area
- responses
- response_functions
- field_responses
- read_field_coordinates

Add context to data: flag to indicate that field coordinates should be read

**Specification**

**Alias:** none

**Argument(s):** none
Field coordinates specify independent variables (e.g. spatial or temporal coordinates) upon which the field depends. For example, the voltage level above might be a function of time, so time is the field coordinate. If the user has field coordinates to read, they need to specify `read_field_coordinates`. The field coordinates will then be read from a file named response_descriptor.coords, where `response_descriptor` is the user-provided descriptor for the field response. The number of columns in the coords file should be equal to the number of field coordinates.

### 7.6.6 no_gradients

- **Keywords Area**
  - responses
  - no_gradients

Gradients will not be used

#### Specification

**Alias:** none  
**Argument(s):** none

**Description**

The `no_gradients` specification means that gradient information is not needed in the study. Therefore, it will neither be retrieved from the simulation nor computed with finite differences. The `no_gradients` keyword is a complete specification for this case.

**See Also**

These keywords may also be of interest:

- `numerical_gradients`
- `analytic_gradients`
- `mixed_gradients`

### 7.6.7 analytic_gradients

- **Keywords Area**
  - responses
  - analytic_gradients

Analysis driver will return gradients

#### Specification

**Alias:** none  
**Argument(s):** none
Description

The `analytic_gradients` specification means that gradient information is available directly from the simulation (finite differencing is not required). The simulation must return the gradient data in the Dakota format (enclosed in single brackets; see Dakota File Data Formats in the Users Manual[5]) for the case of file transfer of data. The `analytic_gradients` keyword is a complete specification for this case.

See Also

These keywords may also be of interest:

- `numerical_gradients`
- `no_gradients`
- `mixed_gradients`

7.6.8 mixed_gradients

- Keywords Area
  - responses
  - mixed_gradients

Gradients are needed and will be obtained from a mix of numerical and analytic sources

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required</td>
<td>id_numerical_-gradients</td>
<td></td>
<td>Identify which numerical gradient corresponds to which response</td>
</tr>
<tr>
<td>Required</td>
<td>id_analytic_-gradients</td>
<td></td>
<td>Identify which analytical gradient corresponds to which response</td>
</tr>
<tr>
<td>Optional</td>
<td>method_source</td>
<td></td>
<td>Specify which finite difference routine is used</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Gradient Source (Group 1)</td>
<td>dakota</td>
<td>(Default) Use internal Dakota finite differences algorithm</td>
</tr>
</tbody>
</table>
Description

The mixed_gradients specification means that some gradient information is available directly from the simulation (analytic) whereas the rest will have to be finite differenced (numerical). This specification allows the user to make use of as much analytic gradient information as is available and then finite difference for the rest.

The method_source, interval_type, and fd_gradient_step_size specifications pertain to those functions listed by the id_numerical_gradients list.

Examples

For example, the objective function may be a simple analytic function of the design variables (e.g., weight) whereas the constraints are nonlinear implicit functions of complex analyses (e.g., maximum stress).

See Also

These keywords may also be of interest:

- numerical_gradients
- no_gradients
- analytic_gradients

7.6.8.1 id_numerical_gradients

- Keywords Area
- responses
- mixed_gradients
- id_numerical_gradients

Identify which numerical gradient corresponds to which response

Topics

This keyword is related to the topics:

- objective_function_pointer
Specification

Alias: none
Argument(s): INTEGERLIST

Description

The \texttt{id\_analytic\_gradients} list specifies by number the functions which have analytic gradients, and the \texttt{id\_numerical\_gradients} list specifies by number the functions which must use numerical gradients. Each function identifier, from 1 through the total number of functions, must appear once and only once within the union of the \texttt{id\_analytic\_gradients} and \texttt{id\_numerical\_gradients} lists.

See Also

These keywords may also be of interest:

- \texttt{id\_analytic\_gradients}

7.6.8.2 \texttt{id\_analytic\_gradients}

- Keywords Area
- responses
- mixed\_gradients
- \texttt{id\_analytic\_gradients}

Identify which analytical gradient corresponds to which response

Topics

This keyword is related to the topics:

- \texttt{objective\_function\_pointer}

Specification

Alias: none
Argument(s): INTEGERLIST

Description

The \texttt{id\_analytic\_gradients} list specifies by number the functions which have analytic gradients, and the \texttt{id\_numerical\_gradients} list specifies by number the functions which must use numerical gradients. Each function identifier, from 1 through the total number of functions, must appear once and only once within the union of the \texttt{id\_analytic\_gradients} and \texttt{id\_numerical\_gradients} lists.

See Also

These keywords may also be of interest:

- \texttt{id\_numerical\_gradients}
7.6.8.3 method_source

- Keywords Area
- responses
- mixed_gradients
- method_source

Specify which finite difference routine is used

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** dakota

**Description**

The `method_source` setting specifies the source of the finite differencing routine that will be used to compute the numerical gradients:

- **dakota** (default)  
- **vendor**

`dakota` denotes Dakota’s internal finite differencing algorithm and `vendor` denotes the finite differencing algorithm supplied by the iterator package in use (DOT, CONMIN, NPSOL, NL2SOL, NLSSOL, ROL, and OPT++ each have their own internal finite differencing routines). The `dakota` routine is the default since it can execute in parallel and exploit the concurrency in finite difference evaluations (see Exploiting Parallelism in the Users Manual [5]). However, the `vendor` setting can be desirable in some cases since certain libraries will modify their algorithm when the finite differencing is performed internally. Since the selection of the `dakota` routine hides the use of finite differencing from the optimizers (the optimizers are configured to accept user-supplied gradients, which some algorithms assume to be of analytic accuracy), the potential exists for the `vendor` setting to trigger the use of an algorithm more optimized for the higher expense and/or lower accuracy of finite-differencing. For example, NPSOL-L uses gradients in its line search when in user-supplied gradient mode (since it assumes they are inexpensive), but uses a value-based line search procedure when internally finite differencing. The use of a value-based line search will often reduce total expense in serial operations. However, in parallel operations, the use of gradients in the NPSOL line search (user-supplied gradient mode) provides excellent load balancing without need to resort to speculative optimization approaches.

In summary, then, the `dakota` routine is preferred for parallel optimization, and the `vendor` routine may be preferred for serial optimization in special cases.

7.6.8.4 dakota

- Keywords Area
- responses
- mixed_gradients
- dakota

(Default) Use internal Dakota finite differences algorithm
Specification

Alias: none
Argument(s): none
Default: relative

Child Keywords:

| Required/Optional Description of Dakota Keyword Dakota Keyword Description |
|-------------------------|---------|---------------------------------|---------------------------------|
| Optional                | Group    | ignore_bounds                   | Do not respect bounds when computing gradients or Hessians (Default) Scale step size by the parameter value |
| Optional/Choose One     | Step Scaling (Group 1) | relative                        |                                   |
|                         |          | absolute                        | Do not scale step-size           |
|                         |          | bounds                          | Scale step-size by the domain of the parameter |

Description

The dakota routine is the default since it can execute in parallel and exploit the concurrency in finite difference evaluations (see Exploiting Parallelism in the Users Manual [5]).

When the method_source is dakota, the user may also specify the type of scaling desired when determining the finite difference step size. The choices are absolute, bounds, and relative. For absolute, the step size will be applied as is. For bounds, it will be scaled by the range of each parameter. For relative, it will be scaled by the parameter value.

7.6.8.4.1 ignore_bounds

- Keywords Area
  - responses
  - mixed_gradients
  - dakota
  - ignore_bounds

Do not respect bounds when computing gradients or Hessians

Specification

Alias: none
Argument(s): none
Default: bounds respected
Description

When Dakota computes gradients or Hessians by finite differences and the variables in question have bounds, it
by default chooses finite-differencing steps that keep the variables within their specified bounds. Older versions
of Dakota generally ignored bounds when computing finite differences. To restore the older behavior, one can add
keyword `ignore_bounds` to the response specification when `method_source dakota` (or just `dakota`) is also specified.

In forward difference or backward difference computations, honoring bounds is straightforward.

To honor bounds when approximating $\frac{\partial f}{\partial x_i}$, i.e., component $i$ of the gradient of $f$, by central differences, Dakota
chooses two steps $h_1$ and $h_2$ with $h_1 \neq h_2$, such that $x + h_1 e_i$ and $x + h_2 e_i$ both satisfy the bounds, and then
computes

$$
\begin{align*}
\frac{\partial f}{\partial x_i} & \approx \frac{h_2^2(f_1 - f_0) - h_1^2(f_2 - f_0)}{h_1 h_2 (h_2 - h_1)},
\end{align*}
$$

with $f_0 = f(x)$, $f_1 = f(x + h_1 e_i)$, and $f_2 = f(x + h_2 e_i)$.

7.6.8.4.2 relative

- Keywords Area
- responses
- mixed_gradients
- dakota
- relative

(Default) Scale step size by the parameter value

Specification

Alias: none

Argument(s): none

Description

Scale step size by the parameter value

7.6.8.4.3 absolute

- Keywords Area
- responses
- mixed_gradients
- dakota
- absolute

Do not scale step-size

Specification

Alias: none

Argument(s): none

Default: relative
7.6 responses

Description
Do not scale step-size

7.6.8.4.4 bounds
  - Keywords Area
  - responses
  - mixed_gradients
  - dakota
  - bounds

Scale step-size by the domain of the parameter

Specification
Alias: none
Argument(s): none

Description
Scale step-size by the domain of the parameter

7.6.8.5 vendor
  - Keywords Area
  - responses
  - mixed_gradients
  - vendor

Use non-Dakota fd algorithm

Specification
Alias: none
Argument(s): none

Description
See parent page for usage notes.

7.6.8.6 interval_type
  - Keywords Area
  - responses
  - mixed_gradients
• interval_type

Specify how to compute gradients and hessians

Specification

Alias: none
Argument(s): none
Default: forward

Description

The interval_type setting is used to select between forward and central differences in the numerical gradient calculations. The dakota, DOT vendor, and OPT++ vendor routines have both forward and central differences available, the CONMIN, NL2SOL and ROL vendor routines support forward differences only, and the NPSOL and NLSSOL vendor routines start with forward differences and automatically switch to central differences as the iteration progresses (the user has no control over this). The following forward difference expression

\[ \nabla f(x) \approx \frac{f(x + he_i) - f(x)}{h} \]

and the following central difference expression

\[ \nabla f(x) \approx \frac{f(x + he_i) - f(x - he_i)}{2h} \]

are used to estimate the \( i^{th} \) component of the gradient vector.

7.6.8.7 forward

• Keywords Area
  • responses
  • mixed_gradients
  • forward

( Default) Use forward differences

Specification

Alias: none
Argument(s): none
Default: forward

Description

See parent page for usage notes.
7.6.8.8 central

- Keywords Area
- responses
- mixed_gradients
- central

Use central differences

Specification

Alias: none
Argument(s): none

Description

See parent page for usage notes.

7.6.8.9 fd_step_size

- Keywords Area
- responses
- mixed_gradients
- fd_step_size

Step size used when computing gradients and Hessians

Specification

Alias: fd_gradient_step_size
Argument(s): REALLIST
Default: 0.001

Description

fd_step_size specifies the relative finite difference step size to be used in the computations. Either a single value may be entered for use with all parameters, or a list of step sizes may be entered, one for each parameter.

The latter option of a list of step sizes is only valid for use with the Dakota finite differencing routine. For Dakota with an interval scaling type of absolute, the differencing interval will be fd_step_size.

For Dakota with and interval scaling type of bounds, the differencing intervals are computed by multiplying fd_step_size with the range of the parameter. For Dakota (with an interval scaling type of relative), DOT, CONMIN, and OPT++, the differencing intervals are computed by multiplying the fd_step_size with the current parameter value. In this case, a minimum absolute differencing interval is needed when the current parameter value is close to zero. This prevents finite difference intervals for the parameter which are too small to distinguish differences in the response quantities being computed. Dakota, DOT, CONMIN, and OPT++ all use \(0.01 \times \text{fd_step_size}\) as their minimum absolute differencing interval. With a \(\text{fd_step_size} = 0.001\), for example,
Dakota, DOT, CONMIN, and OPT++ will use intervals of .001*current value with a minimum interval of 1.e-5. N-PSOL and NLSSOL use a different formula for their finite difference intervals: \( fd\_step\_size \times (1 + \lvert current\_parameter\_value \rvert) \). This definition has the advantage of eliminating the need for a minimum absolute differencing interval since the interval no longer goes to zero as the current parameter value goes to zero.

ROL's finite difference step size can not be controlled via Dakota. Therefore, \( fd\_step\_size \) will be ignored when ROL's finite differencing routines are used (vendor FD gradients are specified). ROL's differencing intervals are computed by multiplying the current parameter value with the square root of machine precision.

### 7.6.9 numerical_gradients

- **Keywords Area**
- **responses**
- **numerical_gradients**

Gradients are needed and will be approximated by finite differences

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td>Gradient Source (Group 1)</td>
<td>method_source</td>
<td>Specify which finite difference routine is used</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Gradient Source (Group 1)</td>
<td>dakota</td>
<td>(Default) Use internal Dakota finite differences algorithm</td>
</tr>
<tr>
<td>Optional (Choose One)</td>
<td>Finite Difference Type (Group 2)</td>
<td>interval_type</td>
<td>Specify how to compute gradients and hessians</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>forward</td>
<td>(Default) Use forward differences</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>central</td>
<td>Use central differences</td>
</tr>
<tr>
<td>Optional</td>
<td></td>
<td>fd_step_size</td>
<td>Step size used when computing gradients and Hessians</td>
</tr>
</tbody>
</table>

**Description**

The `numerical_gradients` specification means that gradient information is needed and will be computed with finite differences using either the native or one of the vendor finite differencing routines.
See Also

These keywords may also be of interest:

- no_gradients
- analytic_gradients
- mixed_gradients

7.6.9.1 method_source

- Keywords Area
- responses
- numerical_gradients
- method_source

Specify which finite difference routine is used

Specification

**Alias:** none

**Argument(s):** none

**Default:** dakota

Description

The **method_source** setting specifies the source of the finite differencing routine that will be used to compute the numerical gradients:

- **dakota** (default)
- **vendor**

**dakota** denotes Dakota’s internal finite differencing algorithm and **vendor** denotes the finite differencing algorithm supplied by the iterator package in use (DOT, CONMIN, NPSOL, NL2SOL, NLSSOL, ROL, and OPT++ each have their own internal finite differencing routines). The **dakota** routine is the default since it can execute in parallel and exploit the concurrency in finite difference evaluations (see Exploiting Parallelism in the Users Manual [5]).

However, the **vendor** setting can be desirable in some cases since certain libraries will modify their algorithm when the finite differencing is performed internally. Since the selection of the **dakota** routine hides the use of finite differencing from the optimizers (the optimizers are configured to accept user-supplied gradients, which some algorithms assume to be of analytic accuracy), the potential exists for the **vendor** setting to trigger the use of an algorithm more optimized for the higher expense and/or lower accuracy of finite-differencing. For example, NPSOL-L uses gradients in its line search when in user-supplied gradient mode (since it assumes they are inexpensive), but uses a value-based line search procedure when internally finite differencing. The use of a value-based line search will often reduce total expense in serial operations. However, in parallel operations, the use of gradients in the NPSOL line search (user-supplied gradient mode) provides excellent load balancing without need to resort to speculative optimization approaches.

In summary, then, the **dakota** routine is preferred for parallel optimization, and the **vendor** routine may be preferred for serial optimization in special cases.
7.6.9.2 dakota

- Keywords Area
- responses
- numerical_gradients
- dakota

(Default) Use internal Dakota finite differences algorithm

**Specification**

**Alias**: none

**Argument(s)**: none

**Default**: relative

**Child Keywords**:

<table>
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<tr>
<th>Optional/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional (Choose One)</td>
<td>Step Scaling (Group 1)</td>
<td>ignore_bounds</td>
<td>Do not respect bounds when computing gradients or Hessians</td>
</tr>
<tr>
<td></td>
<td></td>
<td>relative</td>
<td>(Default) Scale step size by the parameter value</td>
</tr>
<tr>
<td></td>
<td></td>
<td>absolute</td>
<td>Do not scale step-size</td>
</tr>
<tr>
<td></td>
<td></td>
<td>bounds</td>
<td>Scale step-size by the domain of the parameter</td>
</tr>
</tbody>
</table>

**Description**

The `dakota` routine is the default since it can execute in parallel and exploit the concurrency in finite difference evaluations (see Exploiting Parallelism in the Users Manual [5]).

When the `method_source` is `dakota`, the user may also specify the type of scaling desired when determining the finite difference step size. The choices are `absolute`, `bounds`, and `relative`. For `absolute`, the step size will be applied as is. For `bounds`, it will be scaled by the range of each parameter. For `relative`, it will be scaled by the parameter value.

**7.6.9.2.1 ignore_bounds**

- Keywords Area
- responses
- numerical_gradients
- dakota
- ignore_bounds

Do not respect bounds when computing gradients or Hessians
Specification

Alias: none
Argument(s): none
Default: bounds respected

Description

When Dakota computes gradients or Hessians by finite differences and the variables in question have bounds, it by default chooses finite-differencing steps that keep the variables within their specified bounds. Older versions of Dakota generally ignored bounds when computing finite differences. To restore the older behavior, one can add keyword ignore_bounds to the response specification when method_source dakota (or just dakota) is also specified.

In forward difference or backward difference computations, honoring bounds is straightforward.

To honor bounds when approximating \( \frac{\partial f}{\partial x_i} \), i.e., component \( i \) of the gradient of \( f \), by central differences, Dakota chooses two steps \( h_1 \) and \( h_2 \) with \( h_1 \neq h_2 \), such that \( x + h_1 e_i \) and \( x + h_2 e_i \) both satisfy the bounds, and then computes

\[
\frac{\partial f}{\partial x_i} \approx \frac{h_2(f_1 - f_0) - h_1(f_2 - f_0)}{h_1 h_2 (h_2 - h_1)},
\]

with \( f_0 = f(x) \), \( f_1 = f(x + h_1 e_i) \), and \( f_2 = f(x + h_2 e_i) \).

7.6.9.2.2 relative

- Keywords Area
- responses
- numerical_gradients
- dakota
- relative

(Default) Scale step size by the parameter value

Specification

Alias: none
Argument(s): none

Description

Scale step size by the parameter value

7.6.9.2.3 absolute

- Keywords Area
- responses
- numerical_gradients
- dakota
- absolute

Do not scale step-size
Specification

**Alias:** none

**Argument(s):** none

**Default:** relative

**Description**

Do not scale step-size

7.6.9.2.4 bounds

- Keywords Area
- responses
- numerical_gradients
- dakota
- bounds

Scale step-size by the domain of the parameter

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

Scale step-size by the domain of the parameter

7.6.9.3 vendor

- Keywords Area
- responses
- numerical_gradients
- vendor

Use non-Dakota fd algorithm

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

See parent page for usage notes.
7.6.9.4 interval_type

- Keywords Area
- responses
- numerical_gradients
- interval_type

Specify how to compute gradients and hessians

Specification

Alias: none  
Argument(s): none  
Default: forward

Description

The interval_type setting is used to select between forward and central differences in the numerical gradient calculations. The dakota, DOT vendor, and OPT++ vendor routines have both forward and central differences available, the CONMIN, NL2SOL and ROL vendor routines support forward differences only, and the NPSOL and NLSSOL vendor routines start with forward differences and automatically switch to central differences as the iteration progresses (the user has no control over this). The following forward difference expression

$$ \nabla f(x) \approx \frac{f(x + he_i) - f(x)}{h} $$

and the following central difference expression

$$ \nabla f(x) \approx \frac{f(x + he_i) - f(x - he_i)}{2h} $$

are used to estimate the $i^{th}$ component of the gradient vector.

7.6.9.5 forward

- Keywords Area
- responses
- numerical_gradients
- forward

(Default) Use forward differences

Specification

Alias: none  
Argument(s): none  
Default: forward
Description

See parent page for usage notes.

7.6.9.6 central

- Keywords Area
- responses
- numerical_gradients
- central

Use central differences

Specification

Alias: none

Argument(s): none

Description

See parent page for usage notes.

7.6.9.7 fd_step_size

- Keywords Area
- responses
- numerical_gradients
- fd_step_size

Step size used when computing gradients and Hessians

Specification

Alias: fd_gradient_step_size

Argument(s): REALLIST

Default: 0.001

Description

fd_step_size specifies the relative finite difference step size to be used in the computations. Either a single value may be entered for use with all parameters, or a list of step sizes may be entered, one for each parameter.

The latter option of a list of step sizes is only valid for use with the Dakota finite differencing routine. For Dakota with an interval scaling type of absolute, the differencing interval will be fd_step_size.

For Dakota with and interval scaling type of bounds, the differencing intervals are computed by multiplying fd_step_size with the range of the parameter. For Dakota (with an interval scaling type of relative), DOT,
CONMIN, and OPT++, the differencing intervals are computed by multiplying the \texttt{fd\_step\_size} with the current parameter value. In this case, a minimum absolute differencing interval is needed when the current parameter value is close to zero. This prevents finite difference intervals for the parameter which are too small to distinguish differences in the response quantities being computed. Dakota, DOT, CONMIN, and OPT++ all use \texttt{.01*fd\_step\_size} as their minimum absolute differencing interval. With a \texttt{fd\_step\_size = .001}, for example, Dakota, DOT, CONMIN, and OPT++ will use intervals of \texttt{.001*current value} with a minimum interval of \texttt{1.e-5}. NPSOL and NLSSOL use a different formula for their finite difference intervals: \texttt{fd\_step\_size*(1+|current parameter value|)}. This definition has the advantage of eliminating the need for a minimum absolute differencing interval since the interval no longer goes to zero as the current parameter value goes to zero.

ROL's finite difference step size can not be controlled via Dakota. Therefore, \texttt{fd\_step\_size} will be ignored when ROL's finite differencing routines are used (vendor FD gradients are specified). ROL's differencing intervals are computed by multiplying the current parameter value with the square root of machine precision.

7.6.10 no\_hessians

- **Keywords Area**
- **responses**
- **no\_hessians**

Hessians will not be used

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The \texttt{no\_hessians} specification means that the method does not require Dakota to manage the computation of any Hessian information. Therefore, it will neither be retrieved from the simulation nor computed by Dakota. The \texttt{no\_hessians} keyword is a complete specification for this case. Note that, in some cases, Hessian information may still be being approximated internal to an algorithm (e.g., within a quasi-Newton optimizer such as \texttt{optpp\_q\_newton}); however, Dakota has no direct involvement in this process and the responses specification need not include it.

**See Also**

These keywords may also be of interest:

- \texttt{numerical\_hessians}
- \texttt{quasi\_hessians}
- \texttt{analytic\_hessians}
- \texttt{mixed\_hessians}

7.6.11 numerical\_hessians

- **Keywords Area**
- **responses**
• **numerical_hessians**

Hessians are needed and will be approximated by finite differences.

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>fd_step_size</td>
<td>Step size used when computing gradients and Hessians</td>
</tr>
<tr>
<td>Choose One</td>
<td></td>
<td>relative</td>
<td>(Default) Scale step size by the parameter value</td>
</tr>
<tr>
<td></td>
<td>Step Scaling</td>
<td>absolute</td>
<td>Do not scale step size</td>
</tr>
<tr>
<td></td>
<td>(Group 1)</td>
<td>bounds</td>
<td>Scale step-size by the domain of the parameter</td>
</tr>
<tr>
<td></td>
<td></td>
<td>forward</td>
<td>(Default) Use forward differences</td>
</tr>
<tr>
<td></td>
<td></td>
<td>central</td>
<td>Use central differences</td>
</tr>
</tbody>
</table>

**Description**

The **numerical_hessians** specification means that Hessian information is needed and will be computed with finite differences using either first-order gradient differencing (for the cases of **analytic_gradients** or for the functions identified by **id_analytic_gradients** in the case of **mixed_gradients**) or first- or second-order function value differencing (all other gradient specifications). In the former case, the following expression

\[
\nabla^2 f(x)_i \approx \frac{\nabla f(x + he_i) - \nabla f(x)}{h}
\]

estimates the \(i\)th Hessian column, and in the latter case, the following expressions

\[
\nabla^2 f(x)_{i,j} \approx \frac{f(x + he_i + he_j) - f(x + he_i) - f(x + he_j) + f(x)}{h_i h_j}
\]

and

\[
\nabla^2 f(x)_{i,j} \approx \frac{f(x + he_i + he_j) - f(x + he_i - he_j) - f(x - he_i + he_j) + f(x - he_i - he_j)}{4h^2}
\]

provide first- and second-order estimates of the \(ij\)th Hessian term. Prior to Dakota 5.0, Dakota always used second-order estimates. In Dakota 5.0 and newer, the default is to use first-order estimates (which honor bounds on the variables and require only about a quarter as many function evaluations as do the second-order estimates), but specifying **central** after **numerical_hessians** causes Dakota to use the old second-order estimates, which do not honor bounds. In optimization algorithms that use Hessians, there is little reason to use second-order differences in computing Hessian approximations.
See Also

These keywords may also be of interest:

- no_hessians
- quasi_hessians
- analytic_hessians
- mixed_hessians

7.6.11.1 fd_step_size

- Keywords Area
- responses
- numerical_hessians
- fd_step_size

Step size used when computing gradients and Hessians

Specification

**Alias:** fd_hessian_step_size

**Argument(s):** REALLIST

**Default:** 0.001 (forward), 0.002 (central)

Description

`fd_step_size` specifies the relative finite difference step size to be used in the computations. Either a single value may be entered for use with all parameters, or a list of step sizes may be entered, one for each parameter.

The latter option of a list of step sizes is only valid for use with the Dakota finite differencing routine. For Dakota with an interval scaling type of absolute, the differencing interval will be `fd_step_size`.

For Dakota with and interval scaling type of `bounds`, the differencing intervals are computed by multiplying `fd_step_size` with the range of the parameter. For Dakota (with an interval scaling type of `relative`), DOT, CONMIN, and OPT++, the differencing intervals are computed by multiplying the `fd_step_size` with the current parameter value. In this case, a minimum absolute differencing interval is needed when the current parameter value is close to zero. This prevents finite difference intervals for the parameter which are too small to distinguish differences in the response quantities being computed. Dakota, DOT, CONMIN, and OPT++ all use `0.01*fd_step_size` as their minimum absolute differencing interval. With a `fd_step_size = .001`, for example, Dakota, DOT, CONMIN, and OPT++ will use intervals of `.001*current value` with a minimum interval of `1.e-5`. NPSOL and NLSSOL use a different formula for their finite difference intervals: `fd_step_size*(1+|current parameter value|)`. This definition has the advantage of eliminating the need for a minimum absolute differing interval since the interval no longer goes to zero as the current parameter value goes to zero.

ROL's finite difference step size cannot be controlled via Dakota. Therefore, `fd_step_size` will be ignored when ROL's finite differencing routines are used (vendor FD gradients are specified). ROL's differencing intervals are computed by multiplying the current parameter value with the square root of machine precision.
7.6.11.2 relative

- Keywords Area
- responses
- numerical_hessians
- relative

(Default) Scale step size by the parameter value

Specification

Alias: none
Argument(s): none

Description

Scale step size by the parameter value

7.6.11.3 absolute

- Keywords Area
- responses
- numerical_hessians
- absolute

Do not scale step-size

Specification

Alias: none
Argument(s): none
Default: relative

Description

Do not scale step-size

7.6.11.4 bounds

- Keywords Area
- responses
- numerical_hessians
- bounds

Scale step size by the domain of the parameter
7.6 responses

Specification

**Alias:** none

**Argument(s):** none

Description

Scale step-size by the domain of the parameter

7.6.11.5 forward

- **Keywords Area**
- **responses**
- **numerical_hessians**
- **forward**

(Default) Use forward differences

Specification

**Alias:** none

**Argument(s):** none

**Default:** forward

Description

See parent page for usage notes.

7.6.11.6 central

- **Keywords Area**
- **responses**
- **numerical_hessians**
- **central**

Use central differences

Specification

**Alias:** none

**Argument(s):** none

Description

See parent page for usage notes.
7.6.12 quasi_hessians

- Keywords Area
- responses
- quasi_hessians

Hessians are needed and will be approximated by secant updates (BFGS or SR1) from a series of gradient evaluations.

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional Required (Choose One)</th>
<th>Description of Group Quasi-Hessian Approximation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Description</th>
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</thead>
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<tr>
<td></td>
<td></td>
<td>bfgs</td>
<td>Use BFGS method to compute quasi-hessians</td>
</tr>
<tr>
<td></td>
<td></td>
<td>sr1</td>
<td>Use the Symmetric Rank 1 update method to compute quasi-Hessians</td>
</tr>
</tbody>
</table>

Description

The quasi_hessians specification means that Hessian information is needed and will be approximated using secant updates (sometimes called "quasi-Newton updates", though any algorithm that approximates Newton's method is a quasi-Newton method).

Compared to finite difference numerical Hessians, secant approximations do not expend additional function evaluations in estimating all of the second-order information for every point of interest. Rather, they accumulate approximate curvature information over time using the existing gradient evaluations.

The supported secant approximations include the Broyden-Fletcher-Goldfarb-Shanno (BFGS) update (specified with the keyword bfgs) and the Symmetric Rank 1 (SR1) update (specified with the keyword sr1).

See Also

These keywords may also be of interest:

- no_hessians
- numerical_hessians
- analytic_hessians
- mixed_hessians

7.6.12.1 bfgs

- Keywords Area
- responses
Use BFGS method to compute quasi-hessians

Specification

Alias: none
Argument(s): none

Child Keywords:

<table>
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<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>damped</td>
<td>Numerical safeguarding for BFGS updates</td>
</tr>
</tbody>
</table>

Description

Broyden-Fletcher-Goldfarb-Shanno (BFGS) update will be used to compute quasi-Hessians.

\[ B_{k+1} = B_k - \frac{B_k s_k s_k^T B_k y_k y_k^T}{y_k^T s_k} \]

where \( B_k \) is the \( k^{th} \) approximation to the Hessian, \( s_k = x_{k+1} - x_k \) is the step and \( y_k = \nabla f_{k+1} - \nabla f_k \) is the corresponding yield in the gradients.

Notes

- Initial scaling of \( \frac{y_k^T s_k}{y_k^T y_k} \) is used for \( B_0 \) prior to the first update.
- Numerical safeguarding is used to protect against numerically small denominators within the updates.
- This safeguarding skips the update if \( |y_k^T s_k| < 10^{-6} s_k^T B_k s_k \)
- Additional safeguarding can be added using the damped option, which utilizes an alternative damped BFGS update when the curvature condition \( y_k^T s_k > 0 \) is nearly violated.

7.6.12.1.1 damped

- Keywords Area
- responses
- quasi_hessians
- bfgs
- damped

Numerical safeguarding for BFGS updates

Specification

Alias: none
Argument(s): none
Default: undamped BFGS
Use the Symmetric Rank 1 update method to compute quasi-Hessians.

**Specification**

**Alias:** none

**Argument(s):** none

**Description**

The Symmetric Rank 1 (SR1) update (specified with the keyword `sr1`) will be used to compute quasi-Hessians.

\[
B_{k+1} = B_k + \frac{(y_k - B_k s_k)(y_k - B_k s_k)^T}{(y_k - B_k s_k)^T s_k}
\]

where \( B_k \) is the \( k^{th} \) approximation to the Hessian, \( s_k = x_{k+1} - x_k \) is the step and \( y_k = \nabla f_{k+1} - \nabla f_k \) is the corresponding yield in the gradients.

**Notes**

- Initial scaling of \( \frac{y_k^T y_k}{s_k^T s_k} \) is used for \( B_0 \) prior to the first update.
- Numerical safeguarding is used to protect against numerically small denominators within the updates.
- This safeguarding skips the update if \( |(y_k - B_k s_k)^T s_k| < 10^{-6} \|s_k\|_2 \|y_k - B_k s_k\|_2 \)

**7.6.13 analytic_hessians**

**Alias:** none

**Argument(s):** none

Hessians are needed and are available directly from the analysis driver.
Description

The **analytic_hessians** specification means that Hessian information is available directly from the simulation. The simulation must return the Hessian data in the Dakota format (enclosed in double brackets; see Dakota File Data Formats in Users Manual) for the case of file transfer of data. The **analytic_hessians** keyword is a complete specification for this case.

See Also

These keywords may also be of interest:

- **no_hessians**
- **numerical_hessians**
- **quasi_hessians**
- **mixed_hessians**

### 7.6.14 mixed_hessians

- **Keywords Area**
- **responses**
- **mixed_hessians**

Hessians are needed and will be obtained from a mix of numerical, analytic, and "quasi" sources

**Specification**

**Alias:** none

**Argument(s):** none

**Child Keywords:**

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<th>Dakota Keyword Description</th>
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<td>id_numerical_-hessians</td>
<td>Identify which numerical-Hessian corresponds to which response (Default) Scale step size by the parameter value</td>
</tr>
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<td>Optional(Choose One)</td>
<td></td>
<td>relative</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>absolute</td>
<td>Do not scale step-size</td>
</tr>
<tr>
<td></td>
<td>Step Scaling (Group 1)</td>
<td>bounds</td>
<td>Scale step-size by the domain of the parameter</td>
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<th>Finite Difference Type (Group 2)</th>
<th>(Default) Use forward differences</th>
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<tr>
<td>forward</td>
<td></td>
</tr>
<tr>
<td>central</td>
<td></td>
</tr>
</tbody>
</table>

### id_quasi_hessians

Identify which quasi-Hessian corresponds to which response

### id_analytic_hessians

Identify which analytical Hessian corresponds to which response

### fd_hessian_step_size

Identify which numerical-Hessian corresponds to which response

### See Also

These keywords may also be of interest:

- no_hessians
- numerical_hessians
- quasi_hessians
- analytic_hessians

### Topics

This keyword is related to the topics:

- objective_function_pointer
Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

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<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
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<td>Optional</td>
<td></td>
<td>fd_step_size</td>
<td>Step size used when computing gradients and Hessians</td>
</tr>
</tbody>
</table>

**Description**

The `id_analytic_hessians` list specifies by number the functions which have analytic Hessians, and the `id_numerical_hessians` and `id_quasi_hessians` lists specify by number the functions which must use numerical Hessians and secant Hessian updates, respectively. Each function identifier, from 1 through the total number of functions, must appear once and only once within the union of the `id_analytic_hessians`, `id_numerical_hessians`, and `id_quasi_hessians` lists.

**See Also**

These keywords may also be of interest:

- `id_analytic_hessians`
- `id_quasi_hessians`

7.6.14.1.1  fd_step_size

- Keywords Area
- responses
- mixed_hessians
- `id_numerical_hessians`
- `fd_step_size`

Step size used when computing gradients and Hessians

**Specification**

**Alias:** fd_hessian_step_size

**Argument(s):** REALLIST

**Default:** 0.001 (forward), 0.002 (central)

**Description**

`fd_step_size` specifies the relative finite difference step size to be used in the computations. Either a single value may be entered for use with all parameters, or a list of step sizes may be entered, one for each parameter.
The latter option of a list of step sizes is only valid for use with the Dakota finite differencing routine. For Dakota with an interval scaling type of absolute, the differencing interval will be $fd_{\text{step size}}$.

For Dakota with an interval scaling type of bounds, the differencing intervals are computed by multiplying $fd_{\text{step size}}$ with the range of the parameter. For Dakota (with an interval scaling type of relative), DOT, CONMIN, and OPT++, the differencing intervals are computed by multiplying the $fd_{\text{step size}}$ with the current parameter value. In this case, a minimum absolute differencing interval is needed when the current parameter value is close to zero. This prevents finite difference intervals for the parameter which are too small to distinguish differences in the response quantities being computed. Dakota, DOT, CONMIN, and OPT++ all use $0.01 \cdot fd_{\text{step size}}$ as their minimum absolute differencing interval. With a $fd_{\text{step size}} = 0.001$, for example, Dakota, DOT, CONMIN, and OPT++ will use intervals of $0.001 \times \text{current value}$ with a minimum interval of $1.0 \times 10^{-5}$. NPSOL and NLSSOL use a different formula for their finite difference intervals: $fd_{\text{step size}} \times (1 + |\text{current parameter value}|)$. This definition has the advantage of eliminating the need for a minimum absolute differencing interval since the interval no longer goes to zero as the current parameter value goes to zero.

ROL's finite difference step size can not be controlled via Dakota. Therefore, $fd_{\text{step size}}$ will be ignored when ROL's finite differencing routines are used (vendor FD gradients are specified). ROL's differencing intervals are computed by multiplying the current parameter value with the square root of machine precision.

### 7.6.14.2 relative

- **Keywords Area**
  - responses
  - mixed_hessians
  - relative

((Default) Scale step size by the parameter value)

**Specification**

**Alias:** none  
**Argument(s):** none

**Description**

Scale step size by the parameter value

### 7.6.14.3 absolute

- **Keywords Area**
  - responses
  - mixed_hessians
  - absolute

Do not scale step-size

**Specification**

**Alias:** none  
**Argument(s):** none  
**Default:** relative
Description

Do not scale step-size

7.6.14.4 bounds

- Keywords Area
- responses
- mixed_hessians
- bounds

Scale step-size by the domain of the parameter

Specification

Alias: none
Argument(s): none

Description

Scale step-size by the domain of the parameter

7.6.14.5 forward

- Keywords Area
- responses
- mixed_hessians
- forward

(Default) Use forward differences

Specification

Alias: none
Argument(s): none
Default: forward

Description

See parent page for usage notes.
7.6.14.6  central

- Keywords Area
- responses
- mixed_hessians
- central

Use central differences

Specification

**Alias:** none  
**Argument(s):** none

Description

See parent page for usage notes.

7.6.14.7  id_quasi_hessians

- Keywords Area
- responses
- mixed_hessians
- id_quasi_hessians

Identify which quasi-Hessian corresponds to which response

Topics

This keyword is related to the topics:

- objective_function_pointer

Specification

**Alias:** none

**Argument(s):** INTEGERLIST

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group Quasi-Hessian Approximation (Group 1)</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
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<tbody>
<tr>
<td><strong>Required (Choose One)</strong></td>
<td>bfgs</td>
<td>Use BFGS method to compute quasi-hessians</td>
<td></td>
</tr>
</tbody>
</table>
Use the Symmetric Rank 1 update method to compute quasi-Hessians.

Description

The `id_analytic_hessians` list specifies by number the functions which have analytic Hessians, and the `id_numerical_hessians` and `id_quasi_hessians` lists specify by number the functions which must use numerical Hessians and secant Hessian updates, respectively. Each function identifier, from 1 through the total number of functions, must appear once and only once within the union of the `id_analytic_hessians`, `id_numerical_hessians`, and `id_quasi_hessians` lists.

See Also

These keywords may also be of interest:

- `id_numerical_hessians`
- `id_analytic_hessians`

7.6.14.7.1 bfgs

- **Keywords Area**
- responses
- mixed_hessians
- `id_quasi_hessians`
- bfgs

Use BFGS method to compute quasi-hessians

Specification

**Alias:** none

**Argument(s):** none

**Child Keywords:**

<table>
<thead>
<tr>
<th>Required/Optional</th>
<th>Description of Group</th>
<th>Dakota Keyword</th>
<th>Dakota Keyword Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Optional</td>
<td></td>
<td>damped</td>
<td>Numerical safeguarding for BFGS updates</td>
</tr>
</tbody>
</table>

**Description**

Broyden-Fletcher-Goldfarb-Shanno (BFGS) update will be used to compute quasi-Hessians.

\[
B_{k+1} = B_k - \frac{B_k s_k y_k^T B_k}{s_k^T B_k s_k} + \frac{y_k y_k^T}{y_k^T s_k}
\]

where \(B_k\) is the \(t^{th}\) approximation to the Hessian, \(s_k = x_{k+1} - x_k\) is the step and \(y_k = \nabla f_{k+1} - \nabla f_k\) is the corresponding yield in the gradients.

**Notes**

Dakota Version 6.15 Reference Manual generated on November 5, 2021
• Initial scaling of $\frac{y^T_k y_k}{s_k^T s_k}$ is used for $B_0$ prior to the first update.

• Numerical safeguarding is used to protect against numerically small denominators within the updates.

• This safeguarding skips the update if $|y^T_k s_k| < 10^{-6} s_k^T B_k s_k$

• Additional safeguarding can be added using the damped option, which utilizes an alternative damped BFGS update when the curvature condition $y^T_k s_k > 0$ is nearly violated.

7.6.14.7.1.1 damped

• Keywords Area

• responses

• mixed_hessians

• id_quasi_hessians

• bfgs

• damped

Numerical safeguarding for BFGS updates

Specification

Alias: none

Argument(s): none

Default: undamped BFGS

Description

See parent page.

7.6.14.7.2 sr1

• Keywords Area

• responses

• mixed_hessians

• id_quasi_hessians

• sr1

Use the Symmetric Rank 1 update method to compute quasi-Hessians

Specification

Alias: none

Argument(s): none
Description

The Symmetric Rank 1 (SR1) update (specified with the keyword sr1) will be used to compute quasi-Hessians.

\[ B_{k+1} = B_k + \frac{(y_k - B_k s_k)(y_k - B_k s_k)^T}{(y_k - B_k s_k)^T s_k} \]

where \( B_k \) is the \( k \)th approximation to the Hessian, \( s_k = x_{k+1} - x_k \) is the step and \( y_k = \nabla f_{k+1} - \nabla f_k \) is the corresponding yield in the gradients.

Notes

• Initial scaling of \[ \frac{y_k^T y_k}{y_k^T s_k} \] is used for \( B_0 \) prior to the first update.

• Numerical safeguarding is used to protect against numerically small denominators within the updates.

• This safeguarding skips the update if \[ |(y_k - B_k s_k)^T s_k| < 10^{-6}||s_k||_2||y_k - B_k s_k||_2 \]

7.6.14.8 id_analytic_hessians

• Keywords Area
• responses
• mixed_hessians
• id_analytic_hessians

Identify which analytical Hessian corresponds to which response

Topics

This keyword is related to the topics:

• objective_function_pointer

Specification

Alias: none

Argument(s): INTEGERLIST

Description

The id_analytic_hessians list specifies by number the functions which have analytic Hessians, and the id_numerical_hessians and id_quasi_hessians lists specify by number the functions which must use numerical Hessians and secant Hessian updates, respectively. Each function identifier, from 1 through the total number of functions, must appear once and only once within the union of the id_analytic_hessians, id_numerical_hessians, and id_quasi_hessians lists.

See Also

These keywords may also be of interest:

• id_numerical_hessians
• id_quasi_hessians


[77] K. Schittkowski. NLPQLP: A fortran implementation of a sequential quadratic programming algorithm with distributed and non-monotone line search – user’s guide. Technical report, Department of Mathematics, University of Bayreuth, Bayreuth, Germany, 2004. 191


